

MONOGRAPHS ON EXPERIMENTAL BIOLOGY

EDITED BY

JACQUES LOEB, Rockefeller Institute

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VOLUME I

FORCED MOVEMENTS, TROPISMS, AND ANIMAL CONDUCT

BY

JACQUES LOEB, M.D., PH.D., Sc.D.

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PUBLISHED

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FORCED MOVEMENTS, TROPISMS, AND ANIMAL CONDUCT

By JACQUES LOEB, Rockefeller Institute

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FORCED MOVEMENTS, TROPISMS, AND ANIMAL CONDUCT

BY

JACQUES LOEB, M.D., PH.D., Sc.D.

MEMBER OF THE ROCKEFELLER INSTITUTE FOR MEDICAL RESEARCH



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EDITORS' ANNOUNCEMENT

THE rapidly increasing specialization makes it impossible for one author to cover satisfactorily the whole field of modern Biology. This situation, which exists in all the sciences, has induced English authors to issue series of monographs in Biochemistry, Physiology, and Physics. A number of American biologists have decided to provide the same opportunity for the study of Experimental Biology.

Biology, which not long ago was purely descriptive and speculative, has begun to adopt the methods of the exact sciences, recognizing that for permanent progress not only experiments are required but that the experiments should be of a quantitative character. It will be the purpose of this series of monographs to emphasize and further as much as possible this development of Biology.

Experimental Biology and General Physiology are one and the same science, by method as well as by contents, since both aim at explaining life from the physico-chemical constitution of living matter. The series of monographs on Experimental Biology will therefore include the field of traditional General Physiology.

JACQUES LOEB,
T. H. MORGAN,
W. J. V. OSTERHOUT.



AUTHOR'S PREFACE

ANIMAL conduct is known to many through the romantic tales of popularizers, through the descriptive work of biological observers, or through the attempts of vitalists to show the inadequacy of physical laws for the explanation of life. Since none of these contributions are based upon quantitative experiments, they have led only to speculations, which are generally of an anthropomorphic or of a purely verbalistic character. It is the aim of this monograph to show that the subject of animal conduct can be treated by the quantitative methods of the physicist, and that these methods lead to the forced movement or tropism theory of animal conduct, which was proposed by the writer thirty years ago, but which has only recently been carried to some degree of completion. Many of the statements, especially those contained in the first four chapters of the book, are familiar to those who have read the writer's former publications, but so much progress has been made in the last few years that a new and full presentation of the subject seemed desirable. Chapters V to XIII and Chapter XVI are partly or entirely based on new experiments.

Only that part of the literature has been considered which contributes to or prepares the way for quantitative experiments.

The writer is under obligation for valuable criticism to his wife, to Professor T. H. Morgan, and to Lieutenant Leonard B. Loeb, who were kind enough to read the manuscript.

J. L.

The Rockefeller Institute
for Medical Research,
New York.
March, 1918.

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FORCED MOVEMENTS, TROPISMS, AND ANIMAL CONDUCT

CHAPTER I

INTRODUCTION

THE analysis of the mechanism of voluntary and instinctive actions of animals which we propose to undertake in this volume is based on the assumption that all these motions are determined by internal or external forces. Our task is facilitated by the fact that the overwhelming majority of organisms have a bilaterally symmetrical structure, *i.e.*, their body is like our own, divided into a right and left half.

The significance of this symmetrical structure lies in the fact that the morphological plane of symmetry of an animal is also its plane of symmetry in physiological or dynamical respect, inasmuch as under normal conditions the tension in symmetrical muscles is the same, and inasmuch as the chemical constitution and the velocity of chemical reactions are the same for symmetrical elements of the surface of the body, *e.g.*, the sense organs.

Normally the processes inducing locomotion are equal in both halves of the central nervous system, and the tension of the symmetrical muscles being equal, the animal moves in as straight a line as the imperfections of its

locomotor apparatus permit. If, however, the velocity of chemical reactions in one side of the body, *e.g.*, in one eye of an insect, is increased, the physiological symmetry of both sides of the brain and as a consequence the equality of tension of the symmetrical muscles no longer exist. The muscles connected with the more strongly illuminated eye are thrown into a stronger tension,^a and if now impulses for locomotion originate in the central nervous system, they will no longer produce an equal response in the symmetrical muscles, but a stronger one in the muscles turning the head and body of the animal to the source of light. The animal will thus be compelled to change the direction of its motion and to turn to the source of light. As soon as the plane of symmetry goes through the source of light, both eyes receive again equal illumination, the tension (or tonus) of symmetrical muscles becomes equal again, and the impulses for locomotion will now produce equal activity in the symmetrical muscles. As a consequence, the animal will move in a straight line to the source of light until some other asymmetrical disturbance once more changes the direction of motion.

What has been stated for light holds true also if light is replaced by any other form of energy. Motions caused by light or other agencies appear to the layman as expressions of will and purpose on the part of the animal, whereas in reality the animal is forced to go where carried by its legs. For the conduct of animals consists of forced movements.

The term forced movements is borrowed from brain physiology, where it designates the fact that certain animals are no longer able to move in a straight line when

^a We are speaking of positively heliotropic animals exposed to only one source of light.

certain parts of the brain are injured, but are compelled to deviate constantly toward one side, which is (according to the species and the location of the injury in the brain) either the side of the injury or the opposite side. The explanation of these forced movements is that on account of the one-sided injury of the brain the tension of the symmetrical muscles is no longer the same. As a consequence, the impulses for locomotion which are equal for symmetrical muscles will cause greater contraction in certain muscles of one side of the body than in the symmetrical muscles of the other side, and the animal will no longer move in a straight line. The only difference between the forced movements induced by unequal illumination of the two eyes and by injury to the brain is that in the latter case the forced movements may last for days or throughout the whole life, while in the former case they last only as long as the illumination on the two sides of the body is unequal. If we bring about a permanent difference in illumination in the eyes, *e.g.*, by blackening one eye in certain insects, we can also bring about permanent circus motions. This shows that animal conduct may be justly designated as consisting of forced movements.

The idea that the morphological and physiological symmetry conditions in an animal are the key to the understanding of animal conduct demanded that the same principle should explain the conduct of plants, since plants also possess a symmetrical structure. The writer was able to show that sessile animals behave toward light exactly as do sessile plants; and motile animals like motile plants. The forced orientations of plants by outside sources of energy had been called tropisms; and the theory of animal conduct based on the symmetrical struc-

ture of their body was, therefore, designated as the *tropism theory of animal conduct*.

We started with symmetrical animals since in their case the analysis of conduct is comparatively simple; the results obtained in the study of these symmetrical organisms allow us also to understand the conduct of asymmetrical animals. We shall see that the principles underlying their conduct are the same as in the case of symmetrical animals, the asymmetry of the body altering only the geometrical character of the path in which the animal is compelled to move, not, however, the mechanism of conduct. While a perfectly symmetrical organism, possessed of positive heliotropism, moves in a straight line to the source of light, the path deviates from the straight line in the case of an asymmetrical organism and may in some cases, as, *e.g.*, in *Euglena*, be a spiral around the straight line as an axis. Some authors have tried to use asymmetrical organisms as a starting point for the analysis of conduct, but since it is impossible to understand the conduct of the asymmetrical organisms unless it is based upon that of the symmetrical animals, these authors have been led to anthropomorphic speculations, such as "selection of random movements" which, as far as the writer can see, cannot even be expressed in the language of the physicist.

Although the tropism theory of animal conduct was offered thirty years ago ^{285, 286, 287} its acceptance was delayed by various circumstances. In the first place, the majority of the older generation of biologists did not realize that not only the methods of the physicist are needed but also the physicist's general viewpoint concerning the nature of scientific explanation. In many cases the problem of animal conduct is treated in a way

which corresponds more to the viewpoint of the introspective psychologist than to that of the physicist. The attempts to explain animal conduct in terms of "trial and error" or of vague "physiological states" may serve as examples. None of these attempts have led or can lead to any exact quantitative experiments in the sense of the physicist. Other biologists have still more openly adopted an anthropomorphic method of explanation. If pleasure and pain or curiosity play a rôle in human conduct, why should it be otherwise in animal conduct? The answer to this objection is that typical forced movements when produced in human beings, as, *e.g.*, in Ménière's disease or when a galvanic current goes through the brain, are not accompanied by sensations of pleasure or pain, and there is no reason to attribute the circus movements of an animal, after lesion of the brain or when one eye is blackened, to curiosity or thrills of delight. An equally forcible answer lies in the fact that plants show the same tropisms as animals, and it seems somewhat arbitrary to assume that the bending of a plant to the window or the motion of swarmspores of algæ to the window side of a vessel are accompanied or determined by curiosity or by sensations of joy or satisfaction. And finally, since we know nothing of the sentiments and sensations of lower animals, and are still less able to measure them, there is at present no place for them in science.

The second difficulty was created by the fact that the Aristotelian viewpoint still prevails to some extent in biology, namely, that an animal moves only for a purpose, either to seek food or to seek its mate or to undertake something else connected with the preservation of

the individual or the race.^b The Aristotelians had explained the processes in the inanimate world in the same teleological way. Science began when Galileo overthrew this Aristotelian mode of thought and introduced the method of quantitative experiments which leads to mathematical laws free from the metaphysical conception of purpose. The analysis of animal conduct only becomes scientific in so far as it drops the question of purpose and reduces the reactions of animals to quantitative laws. This has been attempted by the tropism theory of animal conduct.

^b This view is still held, especially among authors, who lean more or less openly to vitalism, *e.g.*, v. Uexküll, Jordan, Franz, Bauer, Buddenbrock, and others.

CHAPTER II

THE SYMMETRY RELATIONS OF THE ANIMAL BODY AS THE STARTING POINT FOR THE THEORY OF ANIMAL CONDUCT

THE starting point for a scientific and quantitative analysis of animal conduct is the symmetry relations of the animal body. The existence of these symmetry relations reduces the analysis to a comparatively simple problem.

Organisms show two forms of symmetry, radial symmetry, for which jellyfish and the stems and roots of most plants offer a well-known example, and lateral symmetry, such as exists in man and most animals. In radial symmetry the peripheral elements are distributed equally about an axis of symmetry, in the case of lateral symmetry the peripheral elements are distributed equally to the right and left of the plane of symmetry (or the median plane) by which the body is divided into a right and left half. The importance of this symmetrical structure lies in the fact that the morphological plane of symmetry is also the dynamical plane of symmetry of the organism. Symmetrical spots of the surface of an animal are chemically identical, having the same chemical constitution and also the same quantity of reacting masses. Thus the two eyes are symmetrical organs, each containing the same photochemical substances in equal quantity. In the eye itself each element is to be considered as chemically identical with the symmetrical point in the other eye. Hence, if the two eyes are illuminated equally,

the photochemical reaction products produced in the same time will be equal in both eyes. What is true for the eyes is true for all symmetrical elements of the surface of an animal.

The median plane is also the plane of symmetry for the muscles and the muscular activity of the body. Symmetrical muscles possess under equal conditions equal tension and symmetrical muscles are antagonistic to each other in regard to moving the body to the right or left.

We say that impulses go from the central nervous system to the muscles; and from the surface of the body to the central nervous system. According to our present knowledge that which is called a nervous impulse seems to consist of a wave of chemical reaction traveling along a nerve fiber. The central nervous system is also symmetrical and, moreover, we may conceive a projection of the elements of the surface of the body upon the ganglion cells and from here to the muscular system of the body. The complications in this system of projections constitute the difficulties in our understanding of the structure of the brain. This idea of a projection of the sense organs or the surface of the body upon the brain will explain why the morphological plane of symmetry of an organism is also its plane of symmetry in a dynamical sense. When symmetrical elements of the eyes are struck by light of the same wave length and intensity, the velocity of photochemical reactions will be the same in both eyes. Symmetrical spots of the retina are connected with symmetrical elements in the brain and these in turn are connected with symmetrical muscles. As a consequence of the equal photochemical reactions in the symmetrical spots of the retina equal changes are produced in the symmetrical brain cells with which they are connected, and equal

changes in tension will be produced in the symmetrical muscles on both sides of the body with which the active brain elements are connected.^a On account of the symmetrical character of all the changes no deviation from the original direction of motion will occur. If, however, one eye is illuminated more than the other eye, the influence upon the tension of symmetrical muscles will no longer be the same and the animal will be forced to deviate from the original direction of motion.

We have thus far considered only the relation between right and left. Aside from this symmetry relation we have polarity relations, between apex or head and base or tail end. Just as we found that the morphological plane of symmetry is also a dynamical plane of symmetry, we find that with the morphological polarity head-tail is connected a dynamic polarity of motion, namely, forward and backward. This will become clear in the next chapter on forced movements.

Physiologists have long been in the habit of studying not the reactions of the whole organism but the reactions of isolated segments (the so-called reflexes). While it may seem justifiable to construct the reactions of the

^a Physiologists assume that stimulations are constantly sent from the brain to the muscles and that this maintains their tension. v. Uexküll uses the term that "tonus" is sent out to the muscle and that the brain is a reservoir of "tonus" as if the latter were a liquid. The writer wonders whether it might not be wiser to substitute for such metaphors hypotheses in terms of chemical mass action. Constant illumination causes a stationary process in photosensitive elements of our eye, in which the mass of the reaction product is determined by the Bunsen-Roscoe law. We assume, moreover, that in proportion to this photochemical mass action corresponding chemical reactions take place in the brain elements with which the eyes are connected; and that as a consequence corresponding chemical reactions take place in the muscles by which the tension of the latter is determined. These processes in the muscles may possibly consist in the establishment of a definite hydrogen ion concentration. Such hypotheses have the advantage over the "stimulation" hypothesis that they can be tested by physico-chemical measurements.

organism as a whole from the individual reflexes, such an attempt is in reality doomed to failure, since reactions produced in an isolated element cannot be counted upon to occur when the same element is part of the whole, on account of the mutual inhibitions which the different parts of the organism produce upon each other when in organic connection^b; and it is, therefore, impossible to express the conduct of a whole animal as the algebraic sum of the reflexes of its isolated segments.

E. P. Lyon ³²⁰ has shown that if the tail in a normal shark be bent to one side (without changing the position of the head) the eyes of the animal move as promptly as compass needles in association with the bent tail around the same axis in which the bending occurs, but in an opposite sense. On the convex side of the animal, the white of the eye is more visible in front, on the concave side it is more visible behind; hence the former has moved backward, the latter forward. This was observed not only in the normal fish but also when the optic and auditory nerves were cut. The central nervous system acts as one unit. R. Magnus ³³² and his fellow-workers have shown that an alteration in the position of the head of a dog inevitably alters the tone of the muscles of the legs.^c These and other associations and mutual inhibitions make possible that simplification which allows us to treat the

^b When the stem of a plant (*e.g.*, *Bryophyllum*) is cut into as many pieces as there are nodes, each node will under the proper conditions give rise to one or two shoots. If we leave them in connection, only the buds at the apical end will grow out, the rest of the buds remaining dormant. The whole stem acts as though it consisted of only the bud situated at the apex.

^c The problem of coördination will form the subject of another volume in this series by Professor A. R. Moore, and for this reason a fuller discussion of work on coördination, such as that by Sherrington and by v. Uexküll, may be reserved for Professor Moore's volume.

organism as a whole as a mere symmetry machine, a simplification which forms the foundation of the tropism theory of animal conduct.

It would, therefore, be a misconception to speak of tropisms as of reflexes, since tropisms are reactions of the organism as a whole, while reflexes are reactions of isolated segments. Reflexes and tropisms agree, however, in one respect, inasmuch as both are obviously of a purely physico-chemical character.

CHAPTER III

FORCED MOVEMENTS

WHEN we destroy or injure the brain on one side we paralyze or weaken the muscles connected with this side. As a consequence the morphological plane of symmetry ceases to be the dynamical plane of symmetry and the animal has a tendency to move in circles instead of in a straight line. Suppose a fish swimming forward by motions of its tail fin. Normally the stroke occurs with equal energy to the right and to the left, and the rudder action of the tail is equal in both directions, but after the lesion of one side of the brain the stroke and the rudder action cease to be the same in both directions, it is weakened in one direction. Hence the animal instead of swimming in a straight line is forced to deviate continually toward one side from the straight line of locomotion. We speak in such a case of a forced motion.

When we destroy the ventral portion of the left optic lobe in a shark (*Scyllium canicula*), the fish no longer swims in straight lines but in circles to the right (when the right optic lobe is destroyed it swims in circles to the left). After the destruction of the left optic lobe, the muscles on the left side of the tail are weakened or semi-paralyzed, and they no longer produce the same rudder action as the muscles on the right side. Hence the impulses (or nerve processes) which flow in equal intensity to the muscles on both sides will no longer produce equally energetic rudder action of the tail to the right and to the left, but the muscles turning the tail to the right will

contract more powerfully than those turning it in the opposite direction. The outcome of this greater rudder action of the tail when moving to the right is that the fish instead of swimming in a straight line moves in a circle to the right.²⁹⁰

It is often the case that the body of such a fish even when quiet is no longer straight but bent in a circle, the left side forming the convex side; and when such a fish dies and rigor mortis sets in it may become stiff in this position. These latter observations prove that the circus movements to the right are due to the lowering of the tension of the lateral muscles of the body on the left side of the fish. This is the fundamental fact for the theory of forced movements—namely, that a lesion in one side of the brain lessens the tension of the muscles on one side of the body; as a consequence the motions of the animal become difficult or impossible in one direction and become easy in the opposite direction.

In many cases the motions of an animal depend upon a coöperative activity of two sets of appendages, *e.g.*, the pectoral fins of a fish or the legs of an animal. Such coöperative or associated action is determined by the fact that the same nerve center supplies antagonistic muscles of the two organs (*e.g.*, the lateral fins). Thus the same nerve impulse causes both our eyes to move simultaneously to the right or to the left. When we look to the right, the same impulse which causes the contraction of the rectus externus muscle in the right eye causes a contraction of the rectus internus muscle in the left eye. These two muscles then are associated.

In a fish like the shark the position and innervation of the eyes differ from that of the human being. In the shark the eyes are not in front but on the side, and the

muscles which lift the eye on one side are associated with those which lower it on the other side of the head. A similar association exists in regard to the pectoral fins, the muscles which lift the right pectoral fin are associated with those which lower the left one, and vice versa. When a normal shark swims the two pectoral fins work equally and the fish swims without rolling over to the right or to the left.

If we destroy in a shark the left side of the medulla oblongata forced changes in the position of the two eyes and the two pectoral fins will follow.²⁹⁰ (There are in addition correlated changes in the other fins and the rest of the body which we will omit in order to simplify the presentation of the subject.) When a shark, whose left medulla is cut, is kept in a horizontal position, its left eye looks down and the right eye looks up. This change of position of both eyes indicates that the relative tension between the muscles of the eyes has changed. In the left eye the tension of the lowering muscles predominates over that of their antagonists, in the right eye the reverse is the case. The pectoral fins likewise show associated changes of position. The left fin is raised up dorsally, the right is bent down ventrally. Since we know that the destruction of the central nervous system causes a paralysis of muscles and not the reverse we must conclude that the destruction of the left side of the medulla in a shark causes a weakening or partial paralysis of the muscles which lower the left fin and of those which raise the right fin. Hence the muscles which press down on the water will press harder in the right than in the left fin. When such an animal swims rapidly, it will come under the influence of a couple of forces which must produce a rolling movement around the longitudinal axis of its

body toward the left. These rolling motions are another well-known type of forced movements. When such an animal swims slowly, it will roll more than a normal fish, but it will not roll completely around its longitudinal axis. These are the same motions which were observed in dogs by Magendie and Flourens¹⁵⁵ after an operation in the medulla or pons. We can state, therefore, that the rolling motions are caused by the weakening of one group of (associated) muscles while their antagonists are not weakened.

It is of interest to consider the nature of forced movements after injury of the cerebral hemispheres in a dog. When in a dog one of the cerebral hemispheres is injured the animal immediately after the operation no longer moves in a perfectly straight line, but deviates from the straight line toward that side where the brain is injured.¹⁷⁸ When the left hemisphere is injured circus motions toward the left ensue. If one offers a dog which was operated in the left cerebral hemisphere a piece of meat, removing it as fast as the dog approaches, the dog will move at first a certain distance in a straight line; it will then suddenly turn to the left and describe a complete circle, moving afterward for a little while in a straight line toward the meat and turning again through an angle of 360° to the left, and so on.²³⁴ The explanation is the same as for the foregoing cases. The lesion of the left cerebral hemisphere caused a weakening or partial paralysis of the muscles which turn the body to the right. Hence the animal will, when following the meat, deviate to the left, and this causes a displacement of the retina image in the same direction and an apparent motion of the object to the right. We shall see in a later chapter on

the orienting effect of moving retina images that this deviation of the retina image to the left causes a forced motion of the animal to the right which compensates its tendency to deviate to the left due to the effect of the brain lesion. Hence the animal approaches the meat in an approximately straight line. But it does so with difficulty and sooner or later tiring of this effort it moves in the usual automatic way, whereby equal impulses reach the muscles on both sides. This results in a complete circus movement to the left on account of the weakening (caused by the operation) of muscles which turn the body to the right. The retina image of the meat again induces a straight motion and the whole process described is repeated. When the injury to the brain was less severe the animal may follow the meat for long distances without turning in a circle.

When such a dog is offered simultaneously two pieces of meat, one in front of the left, the other in front of the right eye, it invariably moves toward the one on the left side. The equal flow of impulses caused by the symmetrically located pieces of meat results in a stronger contraction in the muscles on the left than on the right side of the body, since as a consequence of the lesion the tension of the former muscles is greater than that of the latter. When two pieces of meat are simultaneously offered to the dog, but both pieces are in front of the left eye, the dog tries to get the piece nearest to its mouth, but its effort carries it a little too far to the left and then it takes the other piece of meat which is situated farther to the left.²⁸⁴

Some time after the operation these disturbances may become less and may ultimately disappear. If now the

dog is operated on the other, *e.g.*, the right hemisphere, circus motions to the right appear.

We do not wish to exhaust the chapter on forced movements but may perhaps for the sake of completeness point out the following facts. We have seen that if one cerebral hemisphere is injured the dog shows a tendency to circus movements to the operated side. When both hemispheres are injured, *e.g.*, both occipital lobes are removed, the dog can hardly be induced to move forward and it is impossible to cause it to go downstairs, while it is willing to go upstairs. Its front legs are extended and its head is raised high, giving the impression as if such a dog had a tendency to move backward rather than forward or that the forward movement was difficult. If the two anterior halves of the cerebral hemispheres are removed the reverse happens. The animal runs incessantly as if driven by a mad impulse; its head is bent down and it is in every respect the converse of the animal operated in the occipital lobes. These two types of forced movements correspond to the morphological polarity tail-head. This corresponds to the idea of a projection of the surface elements upon the brain either directly or by crossing.

These three types of forced movements: the circus motions, the tendency to go backward, and the irresistible tendency to move forward will appear in the form of the tropistic reactions to be described in this volume.

Since we shall deal in this volume chiefly with invertebrates, it may be of importance to show that forced movements can also be produced in this group of animals by lesion of one side of the cerebral ganglion, and that these forced movements depend also upon the fact that as a consequence of the operation the tension of symmetrical muscles (which is equal under normal condi-

tions) becomes unequal. Fig. 1, *B*, gives the change in position of the body and of the legs in the larva of a dragon fly (*Æschna*) after the left half of the cerebral ganglion has been destroyed (Matula⁵⁴¹). Such an animal moves in a circle to the right. The longitudinal muscles connecting the segments of the body are under higher tension on the right side of the body than on the left and the body

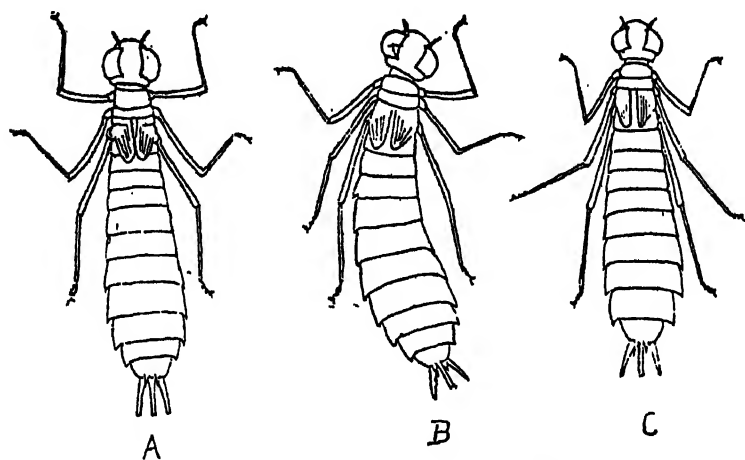


FIG. 1.—*B*, forced position of larva of the dragon fly (*Æschna*) whose left cerebral ganglion is destroyed. The body is convex on the left side, due to a relaxation of the muscles connecting the segments on the left side. The position of the legs is such that the animal can only move in circles to the right. This asymmetry disappears again when both ganglia are destroyed, *C*. *A*, normal animal. (After Matula.)

is bent with its convex side to the left. The normally symmetrical position of the legs (Fig. 1, *A*) is now changed in such a way (Fig. 1, *B*) that the animal is no longer able to move in a straight line, but is forced to move in circles to its right. We shall see later that similar changes in the position of the legs are produced in a positively heliotropic insect when the left eye is blackened and in a negatively heliotropic insect when the right eye

is blackened. Circus motions after destruction of one cephalic ganglion in an insect are a general occurrence and have been known for a long time.

The importance of these forced movements caused by lesion of the brain for the explanation of the conduct of normal animals lies in the fact that the latter is essentially a series of forced movements. The main difference between the forced movements after brain lesion and the conduct of a normal animal lies in the fact that the former are more or less permanent; while in the normal animal conduct the changes in the relative tone of symmetrical muscles leading to a temporary forced movement are caused by a difference in the velocity of chemical reactions in symmetrical sense organs or other elements of the surface.

CHAPTER IV

GALVANOTROPISM

WHEN we send a galvanic current lengthwise through a nerve, at the region near the anode the irritability of the nerve is diminished, while it is increased near the cathode. The condition of decreased irritability near the anode is called anelectrotonus and the increased irritability near the cathode is called catelectrotonus. When a current is sent through an animal, those nerve elements which lie in the direction of the current will have an anelectrotonic and a catelectrotonic region; while the nerves through which the current goes at or nearly at right angles are not affected. Ganglia or nerve tracts in the anelectrotonic condition will, therefore, act as if they were temporarily injured, and hence we need not be surprised to find that the galvanic current causes forced movements which last as long as the current lasts, and which cease with the current.

Hermann reported in 1885²⁰⁴ that when a current is sent through a trough containing tadpoles of a frog, the tadpoles orient themselves in the direction of the current curves putting their heads to the anode.^a Blasius and Schweizer⁵²³ found soon afterwards that a large number of animals when put into a trough with water through which a galvanic current passes have a tendency to go to the anode. The explanation given by Hermann and by Blasius and Schweizer is not correct. They

^a The writer has never been able to repeat this observation.

assumed that the current, acting upon the central nervous system, causes sensations of pain when it goes in the direction from tail to head in the animal; while it has a soothing or hypnotizing effect when it goes in the opposite direction, namely from head to the tail. In the latter case the head is directed toward the anode. The authors assume that the animals choose the position with least pain, *i.e.*, with their heads to the anode. This assumption is wrong, since we know that when a galvanic current is sent through the head of a human being automatic motions comparable to those observed in animals occur which are not voluntary and which are unaccompanied by any pain sensation. Thus when a galvanic current is sent laterally through the head, the person falls toward the anode side but has no feeling of pain. Mach noticed the same effect of falling to the side of the anode when a galvanic current was sent sidewise through fishes.³³⁰ These galvanotropic motions are in reality forced movements, and this has been proved by direct observations. It was shown by Loeb and Maxwell³⁰⁷ in experiments on crustaceans and by Loeb and Garrey³⁰⁸ on salamanders that when we send a galvanic current through animals which go to the anode, changes in the position of the legs occur comparable to the changes in the position of fins and eyes mentioned in the previous chapter, and that these changes are of such a character as to make it easy for the animal to move in the direction of the anode and difficult if not impossible to move in the opposite direction.

In all these experiments it is of importance to choose the proper density of the current. For the experiments on the shrimp (*Palæmonetes*)³⁰⁷ the animals were put into a

square trough, two opposite sides of which were formed of platinum electrodes. The cross section of the *fresh* water in the trough was 1,400 mm.² and the intensity of the current about 1 milliampere or a little less. We found it advisable to increase the intensity very gradually by increasing slowly the resistance of a rheostat in a short circuit until the phenomenon of galvanotropism appeared most strikingly. When the current is too strong or too weak the phenomena are no longer clear. The common shrimp (*Palæmonetes*) is a marine crustacean which

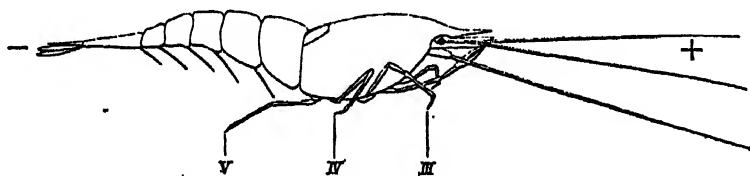


FIG. 2.—Forced position of shrimp (*Palæmonetes*) when galvanic current goes from head to tail. Tension of extensor muscles of tail fin prevails over that of flexors. Animal can swim forward (to anode), but not backward. (After Loeb and Maxwell.)

lives also in brackish water and which can stand fresh water long enough for the purpose of these experiments. The animal can swim forward as well as backward; in forward swimming the extensor muscles of its tail fin work more strongly than the flexors (Fig. 2); in swimming backward the flexors work energetically (Fig. 3) and thus produce a powerful stroke forward, while the extensors contract with less energy. When we put a *Palæmonetes* in a trough through which a current goes and if we put the animal with its head toward the anode the tail is stretched out (Fig. 2). This means that the tension of the extensor muscles prevails over that of the flexors and since the forward swimming is due to the stroke of

the extensors, and since it is antagonized by the tension of the flexors, the animal can swim forward but not backward, or only with difficulty; if we put the animal with its head toward the cathode the tail is bent ventrally (Fig. 3), which means that the tension of the flexors is stronger than that of the extensors. As a consequence the animal can swim backward but not forward, or only

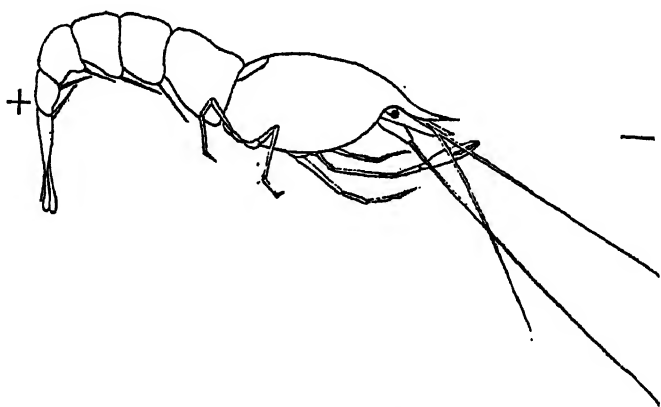


FIG. 3.—Forced position of shrimp when positive current goes from tail to head. Tension of flexors of tail fin prevails over that of extensors. Animal can swim backward (to anode), but not forward. (After Loeb and Maxwell.)

with difficulty. In both cases the result will be a swimming of the animal to the anode, in the former case by swimming forward in the latter by swimming backward.

We can further show that the tension of the muscles of the legs of *Palamonetes* is always altered in such a sense by the galvanic current that motion toward the anode is facilitated, while that toward the cathode is rendered difficult or impossible. The animal uses the third, fourth, and fifth pair of legs for its locomotion (Fig. 2). The third pair pulls in the forward movement

and the fifth pair pushes. The fourth pair acts like the fifth and requires no special discussion. If a current be sent through the animal longitudinally from head to tail and the intensity be increased gradually, a change soon takes place in the position of the legs. In the third pair the tension of the flexors predominates (Fig. 2), in the fifth the tension of the extensors. The animal can thus move easily by pulling of the third and by pushing of the fifth pair of legs, that is to say, the current changes the tension of the muscles in such a way that the forward motion is facilitated, while the backward motion is rendered difficult. Hence it can easily go toward the anode but only with difficulty toward the cathode. If a current be sent through the animal in the opposite direction, namely from tail to head, the third pair of legs is extended, the fifth pair bent (Fig. 3); *i.e.*, the third pair can push, the fifth pair can pull backward. The animal can thus go backward with ease but forward only with difficulty. This again will lead to a gathering of such animals at the anode, this time, however, by walking backward.

The phenomena thus far described recall the forced movements mentioned in the third chapter, where certain injuries of the brain accelerate forward motion while other lesions in the opposite parts of the brain make forward motion difficult if not impossible.

Palæmonetes can also walk sidewise. This movement is produced by the pulling of the legs on the side toward which the animal is moving (contraction of the flexors), while the legs of the other side push (contraction of extensors). If a current be sent transversely, say from left to right, through the animal, the legs of the left side assume the flexor position, those of the right side the

extensor position (Fig. 4). The transverse current thus makes it easy for the animal to move toward the left—the anode—and prevents it from moving toward the right—the cathode. If a galvanic current flows transversely

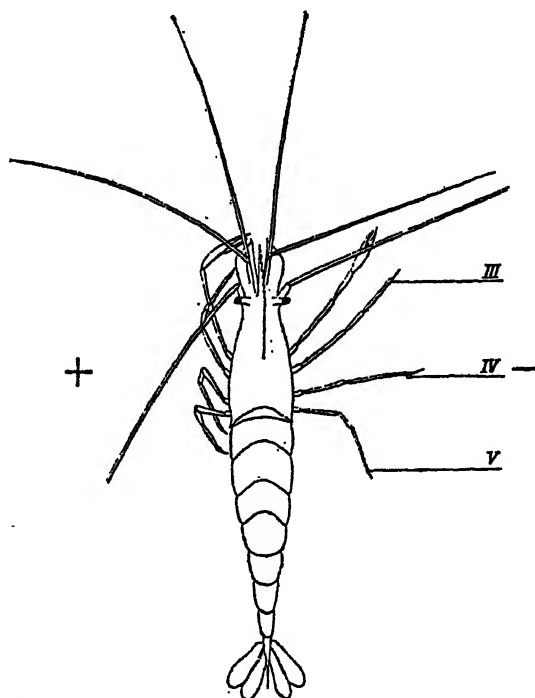


FIG. 4.—Position of legs of shrimp when current goes sidewise through the animal, from left to right. In the legs on the left side the tension of the flexors, in those of the right side the tension of the extensors predominates. The animal can easily go to the left (anode), but not to the right. (After Loeb and Maxwell.)

through the animal, it creates the analogue of the circus motions produced by injury of one side of the brain. Figs. 5 and 6 show that the current produces similar effects in the crayfish as those produced in the shrimp (Figs. 2 and 3).

It is not difficult to suggest by aid of a diagram the arrangement of the elements in the central nervous system required to bring about the phenomena of galvanotropism just described for *Palæmonetes*. We take it for granted that the regular phenomena of anelectrotonus and catelectrotonus of motor nerve elements suffice for the explanation of these phenomena. We assume that if the cell body of a neuron is in the state of catelectrotonus

FIG. 5.

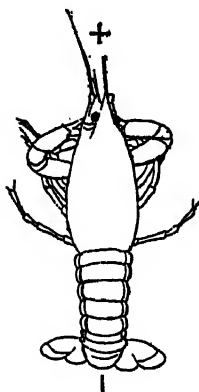
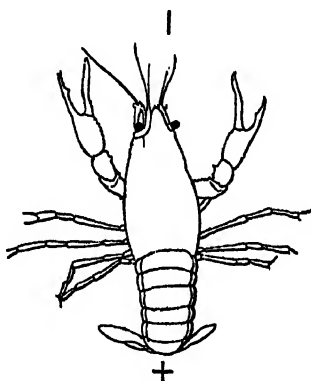


FIG. 6.



FIGS. 5 and 6.—Show the same effects of current on the common crayfish as those on *Palæmonetes* in Figs. 2 and 3.

its activity is increased, when it is in anelectrotonic condition activity is diminished. Neurons having the same orientation will always be affected in the same sense by the current.

Fig. 7 is a diagram of the arrangement of neurons giving rise to the bending of the legs on the side of the anode and to the extension of the legs on the side of the cathode when the current goes sidewise through the animal. This diagram assumes that the nerves innervating the extensors come from the opposite side of the central

nervous system, while those innervating the flexors are on the same side. This diagram corresponds to reality, according to the histological work of Allen. When the current goes from right to left through the crustacean the cell bodies of the neurons on the right side are in catelectrotonus, those on the left side in anelectrotonus. The former are, therefore, in a state of increased "irritability," the latter in a state of diminished "irritability." Hence the flexors of the right leg are contracted and the extensors relaxed, while the flexors of the left leg are relaxed and the extensors contracted.

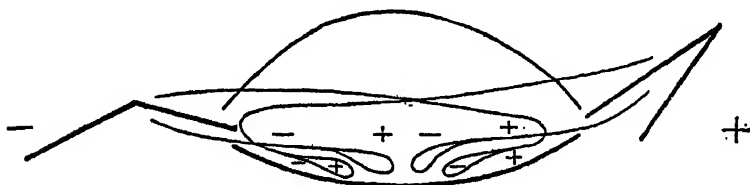


FIG. 7.—Diagram indicating the orientation of the neurons for flexor and extensor muscles of the right and left legs to explain changes of position of legs under influence of galvanic current. (After Loeb and Maxwell.)

Another crustacean *Gelasimus*³⁰⁷ shows the same effect of the current when it goes sidewise through its body. When the thoracic ganglion from which the nerves of the legs originate is cut longitudinally in the middle, all the legs assume permanently a bent position, confirming our assumption that the extensor nerves cross over while the flexors originate from the same side of the ganglion on which their muscles are. It, therefore, looks as if our diagram were the expression of the actual condition.

In the same way we can explain the results of a galvanic current when it goes through the animal lengthwise. We only need to assume that the cell bodies which send their fibers to the flexors of the third pair of legs

have the same orientation as the cell bodies which send their fibers to the extensors of the fifth pair of legs (Fig. 8). Hence when the positive current goes from head to tail through the animal (Fig. 8), the flexors of the third pair of legs and the extensors of the fifth pair must be thrown into greater activity, since the cell bodies of both these nerves are in a condition of catelectrotonus, *i.e.*, increased activity.

FIG. 8.

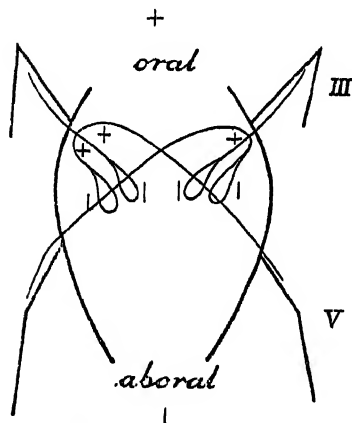
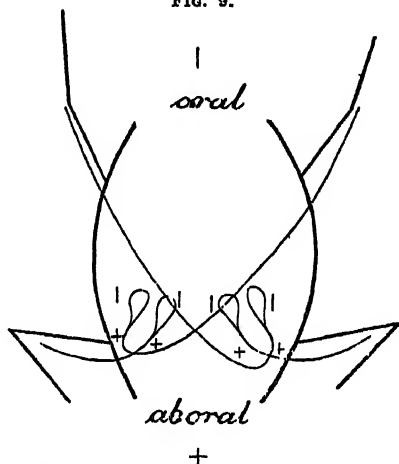


FIG. 9.



FIGS. 8 and 9.—Diagram indicating orientation of neurons for flexor and extensor muscles of third and fifth pairs of legs to explain galvanotropic reaction. (After Loeb and Maxwell.)

When the current goes from tail to head the cell bodies of the extensors of the third and of the flexors of the fifth pair of legs are in catelectrotonus. This possibility is expressed in the diagram Fig. 9.

In this way the theory of the galvanotropic reaction of those animals which go to the anode seems complete.

What has been demonstrated for *Palæmonetes* holds not only for many crustaceans but for vertebrates also. Loeb and Garrey³⁰⁶ have shown that when a current

is sent through a trough containing larvæ of a salamander (*Amblystoma*) the legs and head of the larvæ assume definite positions depending upon the direction of the current. When the current goes from head to tail the legs are pushed backward and the head is bent (Fig. 10);



FIG. 10.—Forced position of *Amblystoma* larva under influence of galvanic current going through animal from head to tail. Head down, body convex on dorsal side. Legs backward. (After Loeb and Garrey.)

when the current goes from tail to head the opposite position is observed (Fig. 11). The analogy with the observations on *Palæmonetes* is obvious.

Galvanotropic reactions are found throughout the whole animal kingdom and the following observations made by Bancroft on a jellyfish (*Polyorchis penicillata*)

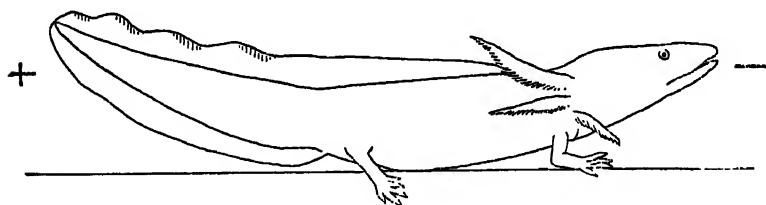


FIG. 11.—Forced position of *Amblystoma* larva when current goes from tail to head. Head raised, legs pushed forward, tail raised. (After Loeb and Garrey.)

are of especial interest.¹⁸ Strips containing tentacles and the manubrium were cut out from the animal and put into a trough through which a current flowed of 25 to 0.200 m. a. for 1 square mm. of the cross section of the liquid (dilute sea water) in the trough.

If a meridional strip passing from the edge on one side through the center of the bell to the other edge be prepared and the current passed through transversely, tentacles and manubrium turn and point toward the cathode (Fig. 12). A reversal of the current initiates a turning of these organs in the opposite direction, which is usually completed in a few seconds. This can be repeated many times and the tentacles continue to respond after hours of activity. The manubrium,

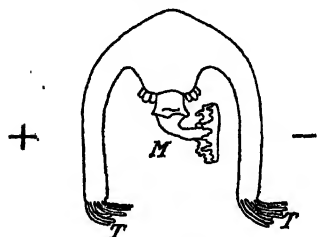


FIG. 12.—Tentacles *T* and manubrium *M* of a jellyfish (*Polyorchia*) under influence of galvanic current are turned to the negative pole. (After Bancroft.)

however, tires sooner and fails to respond. If the strip is placed with its subumbrella surface upward and extended in a straight line parallel to the current lines, the making of the current causes the tentacles at the anode end to turn through an angle of 180° and point toward the cathode. The tentacles at the cathode end become more crowded together, reminding one of the tip of a moistened paint brush, and also point more directly toward the cathode (Fig. 13). The experiment may be varied in still other ways by

cutting smaller or larger pieces from the edge of the swimming bell, but the response is always the same. The tentacles, wherever possible, and to a less extent the manubrium, bend so as to point toward the cathode. The response depends in no way upon the connection of these organs with the swimming bell, muscles, or nerve ring, for it is obtained equally well with isolated tentacles and pieces of tentacles. Isolated tentacles when placed transversely to



FIG. 13.—Strip of jellyfish showing that under the influence of galvanic current tentacles on both ends point towards cathode. (After Bancroft.)

the current lines curve so as to assume a more or less complete U-shape, with their concave side toward the cathode. When placed parallel to the current, the tentacles do not curve.²⁰

The latter observation shows the fact that the whole reaction is due merely to an increase in the tension of the muscles on the cathode side of the organ.

Phenomena of galvanotropism can be observed also in infusorians. Thus Verworn⁴⁹³ observed that when

a current goes through a trough containing *Paramæcia* the animals will all move toward the cathode. The mechanism of the reaction was discovered by Ludloff.³¹⁷ The locomotion of *Paramæcium* is brought about by cilia. As a rule the cilia are directed backward (Fig. 14), and in their normal movement they strike powerfully backward and are retracted with less energy to their normal position. Since their powerful stroke is backward the animal is pushed forward in the water. Ludloff and Bancroft^{17, 18} show that if a *Paramæcium* is struck sideways by the current, the position of the cilia on the cathode side is reversed inasmuch as they are now turned forward. On the anode side they continue to be directed backward (Fig. 15, *a*). Instead of striking symmetrically on both sides of the animal, the cilia on the cathode side strike forward powerfully while those on the anode side strike powerfully backward. The animal is thus under the influence of a couple of forces which turn its oral pole toward the cathode side. As soon as it is in this condition the symmetrical cilia are struck at the same angle by the parallel current lines and they must assume a symmetrical position which is as in Fig. 15, *b*, namely the cilia are pointed forward toward the cathode at the oral end, and backward toward the anode at the aboral end. As long as the current is not too strong, the oral region, where the cilia are pointing forward, is rather small and therefore the action of those cilia prevails which are in the majority and which are pointed backward. As a result the organism moves slowly forward to the cathode.



FIG. 14.—*Paramæcium* under normal conditions. Cilia all pointing toward aboral pole.

A similar mechanism of galvanotropic conduct exists in *Volvox* a spherical, unicellular organism which is surrounded by cilia on its whole surface. A definite pole of the organism is always foremost in all locomotions. This organism usually swims to the anode when in a galvanic field. Bancroft made the action of the cilia of *Volvox* visible with the aid of india ink and was able to show that the current made the cilia on the anode side stop, while those on the cathode side continue to beat.²⁰

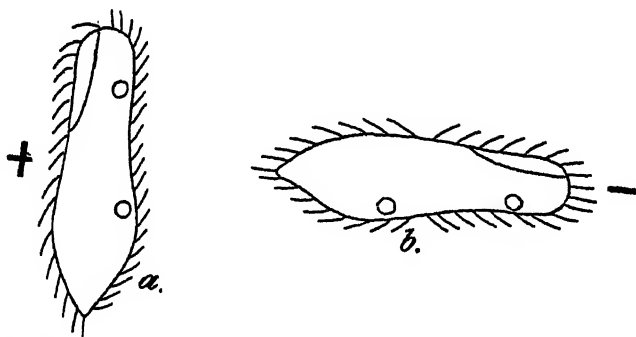


FIG. 15.—*a*, current going from left to right through *Paramaecium*, the position of cilia on the cathode side is now reversed, their free ends pointing forward. The animal when swimming is automatically turned with its oral end toward the cathode. *b*, current going through *Paramaecium* from aboral to oral end. Cilia symmetrical on both sides but pointing forward at oral end and backward at aboral end.

Since the backward stroke is always the effective one the organism is thus carried automatically toward the anode.

Terry⁴⁷⁸ found that *Volvox* can be made to move toward the anode as well as toward the cathode. It moves to the anode after having been kept in the dark for two or three days, while after exposure to light it swims to the cathode. *Volvox* contains chlorophyll and the change in the sense of reaction is therefore connected with the formation of a product of chlorophyll activity. Bancroft found that when *Volvox* was made cathodic by exposure to sunlight, the cilia stop on the cathode side.

While the locomotor mechanism of unicellular organisms, like *Paramæcia* and *Volvox*, is as simple as that of higher organisms, the locomotion of microörganisms possessing only one flagellum, like *Euglena*, is more complicated. It was generally assumed that the flagellum acted like a single oar and that it was directed forward, but this is not correct. It is shaped like a U and its free end is directed backward; and Bancroft has emphasized that it acts by the formation of a loop which moves like a wave from the base of the flagellum to its free tip. The same author discovered that *Euglena* are galvanotropic when raised in acid media. On account of the asymmetry of their locomotor apparatus they are compelled to swim in a spiral, in most cases to the cathode, exceptionally to the anode. Bancroft showed that the orientation of these organisms by the galvanic current is identical with that by light.²¹

All the phenomena of galvanotropism are, therefore, reduced to changes in the tension of associated muscles or contractile elements, as a consequence of which the motion of the organism toward one pole is facilitated, while the motion toward the opposite pole is rendered difficult. Galvanotropism is, therefore, a form of forced motions produced by the galvanic current instead of by injury to the brain.

There remains then the question of how a galvanic current can bring about those changes which result in the anelectrotonic and catelectrotonic condition mentioned at the beginning. Currents can pass through tissues only in the form of ions whose progress is blocked by membranes which are more permeable for certain salts than for others. Those salts which go through the membrane carry the current through the tissue elements, those

which do not go through will increase in concentration at the surface of the membrane. It is the latter which cause the electrotonic effects; according to Loeb and Budgett³⁰⁴ by secondary chemical reactions at the boundary. Nernst has pointed out that a stationary condition must arise at the surface of the membrane due to the fact that the increase in concentration of ions by the electric current gives rise to a current of diffusion of salt in the opposite direction away from the membrane. "The average change of concentration at the membrane depends, therefore, upon the antagonistic effects of the current and of the diffusion."⁵²⁴ This must be kept in mind since otherwise the effect of the constant current should increase constantly with its duration, which is not the case, on account of the establishment of a condition of equilibrium between the increase of the concentration of ions at the boundary with the duration of the current and the diminution of this concentration by the diffusion of the ions in the opposite direction due to osmotic pressure.

CHAPTER V

HELIOTROPISM

THE INFLUENCE OF ONE SOURCE OF LIGHT

1. GENERAL FACTS

THE fact that certain animals go to the light had, of course, been known for hundreds of years, but this was explained in an anthropomorphic way. Thus Lubbock, and Graber,¹⁸⁰ had taken it for granted that certain animals went to the light or away from it on account of fondness for either light or darkness, and their experiments were calculated to demonstrate this fondness. Animals were distributed in a box, one-half of which was covered with common window glass, the other with an opaque body or with colored glass, and after a while the number of animals in each half was counted. When the majority of animals were found in the dark part the animals were believed to have a preference for darkness, when in the light part they were believed to be fond of light. The same method was used to decide whether animals preferred blue to red or vice versa. The writer attacked the problem from the physical viewpoint, assuming that the animals are "fond" neither of light nor of "darkness," but that they are oriented by the light in a similar way as plants are; being compelled to bend or—as in the case of motile algæ—move automatically either to a source of light or away from it.^{285, 287}

In the case of unequal illumination of the two eyes the tension of the symmetrical muscles in an animal becomes

unequal. In this condition the equal impulses of locomotion will result in an unequal contraction of the muscles on both sides of the animal. As a consequence the animal will turn automatically until its plane of symmetry is in the direction of the rays of light. As soon as this happens the illumination of both eyes and the tension of symmetrical muscles become equal again and the animal will now move in a straight line—either to or from the source of light. What appeared to the older authors as the expression of fondness for light or for darkness was according to the writer's theory the expression of an influence of light upon the relative tension of symmetrical muscles.

Animals which are compelled to turn and move to the source of light the writer called positively heliotropic, those which are compelled to turn and move in the opposite direction he called negatively heliotropic. The designation heliotropism (or phototropism) was chosen to indicate that these reactions of animals are of the same nature as the turning of plants to the light; and the writer was indeed able to show that sessile animals bend to the light as do plants which are raised near a window;²⁸⁸ while motile animals move to (or from) a source of light as do the motile swarmspores of algæ or motile algæ themselves.

We will first discuss positively heliotropic motile animals. The positively heliotropic caterpillars of *Porthesia chrysorrhæa*²⁸⁸ or the winged plant lice of *Cineraria*²⁸⁸ or the newly hatched larvæ of the barnacle¹⁸³ were used by the writer in his earliest experiments and they may serve as examples. The larvæ of *Porthesia* must be used after hibernation before they have taken food. When about 50 or 100 of such larvæ are put into a test tube and

the latter is placed at right angles against a window, all the animals begin to move to the window in as straight a line as the imperfections of their locomotion and collisions permit. As soon as they reach the window side of the test tube they remain there permanently, unless the test tube is turned around. If we turn the test tube around an angle of 180° the animals go at once to the window again. They react in this way whether the source of light is sunlight, diffused daylight, or lamp light; and this can be repeated indefinitely. The animals are slaves of the light. These experiments are typical for positively heliotropic motile animals.

When the animals have reached the window end of the test tubes they remain there, since the light prevents them from going back. But in staying there they may assume any kind of orientation, thus proving that the light orients them only as long as they are in motion. The light affects the tension of the muscles and we shall see later that when the animals are not moving, the change in the tension of the muscles manifests itself by changes in the position of the legs, which is noticeable in organisms with comparatively large appendages.

That these animals do not go to the light because they prefer light to darkness but because the light orients them is proved by the fact that they will go from light into the shade if by so doing they remain oriented with their heads toward the source of light.²⁸⁷ Let direct sunlight S fall upon a table through the upper half of a window (W , Fig. 16), the diffused daylight D through the lower half. A test tube ac is placed on the table in such a way that its long axis is at right angles with the plane of the window; and one-half ab is in the direct sunlight, the other half in the shade. If at the beginning of the

experiment the positively heliotropic animals are in the direct sunlight at *a*, they promptly move toward the window, gathering at the window end *c* of the tube, although by so doing they go from the sunshine into the shade. This experiment shows also that it is not the intensity

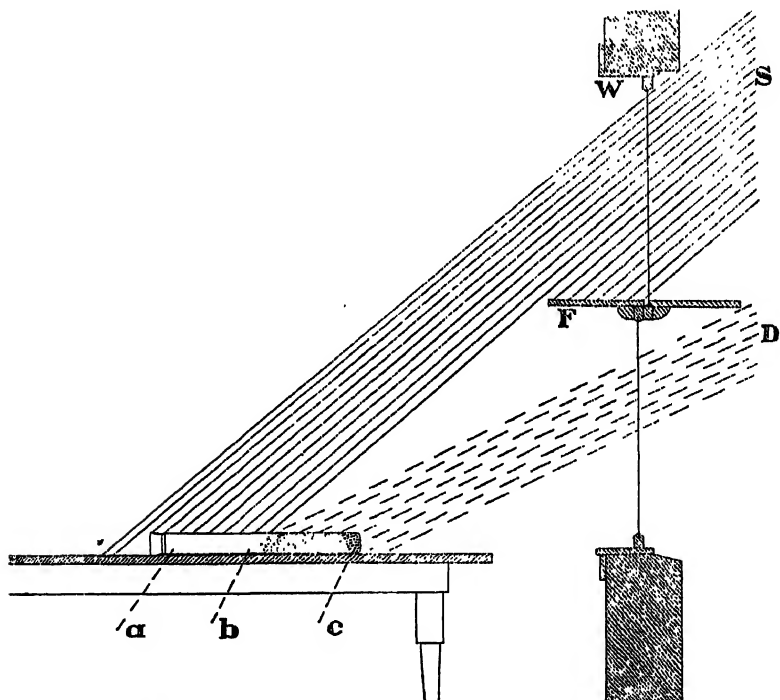


FIG. 16.—Showing that positively heliotropic animals will move from sunlight into shade if in so doing the illumination of the two eyes remains the same.

gradient of light in the dish which makes positively heliotropic animals move to the light, but that difference in intensity on both sides of the animal which is caused by the screening effect of the animal's own body. The same holds true for chemotropism.

Thus far we have discussed positively heliotropic

animals only. In the case of unequal illumination of the two eyes or of the two sides of the body of a negatively heliotropic animal the tension in the muscles turning the animal to the source of light is diminished. The impulses for locomotion which are equal for the muscles of both sides of the body will, therefore, result in turning the head of the animal away from the source of light. As soon as the plane of symmetry of the animal goes again through the source of light, the symmetrical photosensitive elements of the head receive again equal illumination, and the animal will now continue to move in a straight line away from the source of light. The fully grown larvæ of the housefly when they are ready to pupate show this negative heliotropism.

Negatively heliotropic animals, *e.g.*, the fully grown larvæ of the blowfly, can be made to move from weak light to strong light, *e.g.*, from the shade into direct sunlight, if in so doing the illumination on the two sides of the body remains equal.²⁸⁷ This was shown by the writer by an arrangement similar in principle to the one described above. Thus the idea that the intensity gradient of light determines the direction of motion was disproved also for negatively heliotropic animals.

Thus far we have shown only that a heliotropic animal is oriented in such a way to a source of light that its plane of symmetry goes through the source of light. This does not yet explain why a positively heliotropic animal cannot go away from the source of light, since in going to or going away from the source of light both sides of the animal receive equal illumination. The fact that a positively heliotropic animal cannot go away from the light finds its explanation by observations of Holmes²²⁸ and Garrey,¹⁷⁷ showing that when light falls from behind

and above on a positively heliotropic animal its progressive motions are stopped, and in some cases a tendency to turn a somersault backward may even arise. The case is similar to that of galvanotropism when the current goes through an animal lengthwise (see previous chapter). We must conclude from the observations of Holmes and Garrey, which will be discussed farther on, that if the head of a positively heliotropic animal is turned to a source of light its forward motions are facilitated and the backward motions rendered difficult; while in the case of a negatively heliotropic animal it is just the reverse. If the animal now moves to the right or to the left the illumination of the two eyes or of the two sides of the body becomes different again, causing a forced movement, whereby the plane of symmetry of the moving animal is caused to go through the source of light again; with the head toward the source of light when the animal is positively heliotropic or away from it when it is negatively heliotropic.

2. DIRECT PROOF OF THE MUSCLE TENSION THEORY OF HELIOTROPISM IN MOTILE ANIMALS

The fact that light causes forced movements, like those described in the case of brain lesions and of galvanotropism, has been proved by many observers, and especially clearly by Holmes and Garrey. Holmes worked on the positively heliotropic water scorpion *Ranatra*.²²⁸ When this animal is illuminated from the right side, the legs on the right side of the body are bent and those on the left side extended (Fig. 17). This effect is identical with the one observed in *Palæmonetes*, when a galvanic current goes sidewise through the animal. Hence *Ranatra*

can easily move to the source of light on its right side but with difficulty or not at all in the opposite direction.

When the light is placed behind the animal, the body is raised up in front and the head held high in the air (Fig. 18). The opposite attitude is assumed, when the light is placed in front, the body being lowered and the head bent down (Fig. 18). These effects resemble the

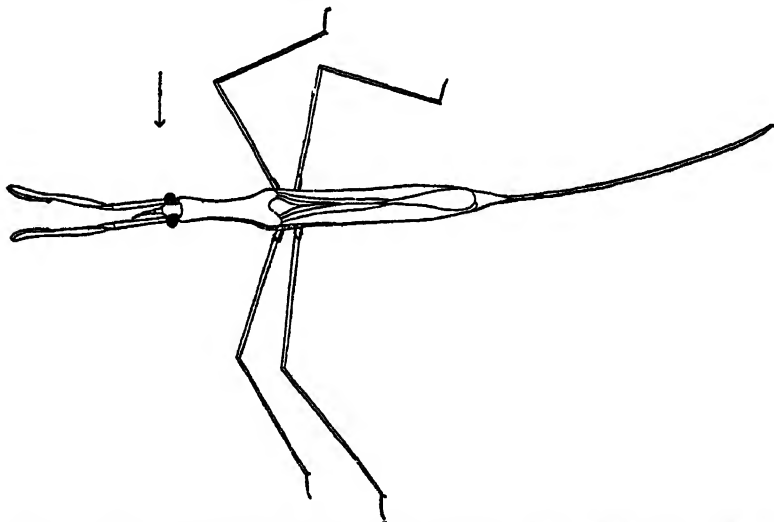


FIG. 17.—Position of the water scorpion *Ranatra* when the right eye is toward the light. (After Holmes.)

galvanotropic effects observed in the position of the head of *Amblystoma* when the current goes forward or backward through the animal.

These latter observations of Holmes explain, as already mentioned, why a positively heliotropic animal cannot move away from the light and why a negatively heliotropic animal cannot move to a source of light. The progressive motions of the negatively heliotropic animal will be stopped when the light strikes it in front; while

these motions of the positively heliotropic animal will be facilitated when the light is in front and will be rendered impossible when the light is behind.

The writer had observed long ago that when the convexity of one eye is cut off in the housefly it will no longer go in a straight line but will make circus movements, the normal eye being directed toward the center of the circle.²⁸⁶ It was shown by Parker that blackening of one

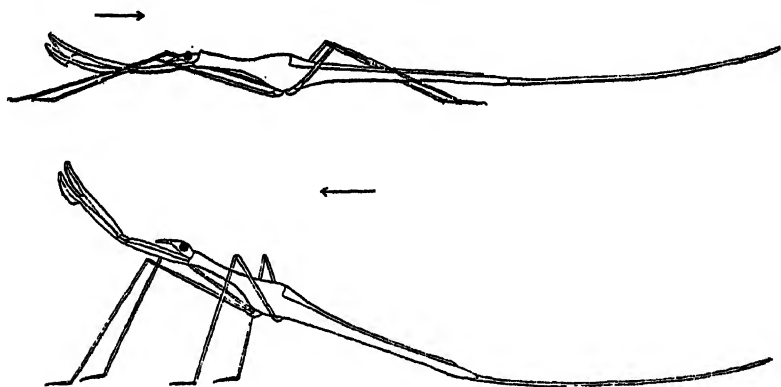


FIG. 18.—The lower figure represents the position of *Ranatra* when the light is behind the body. The upper figure represents the position assumed when the light is in front of the animal. (After Holmes.)

eye of the positively heliotropic butterfly *Vanessa antiopa* calls forth circus movements, with the unblackened eye toward the center of the circle.³⁹⁸ Holmes, Rádl,⁴⁴⁷ Axenfeld, Garrey,¹⁷⁷ and many other authors have since made similar observations. In the positively heliotropic *Ranatra*, Holmes described the effect of blackening one eye as follows:

If one eye of *Ranatra* is blackened over or destroyed the insect in most cases no longer walks in a straight line but performs more or less decided circus movements toward the normal side. Under the stimulus of light the insect assumes a peculiar attitude; the body leans over toward the normal side and the head is tilted over in the same direction.²²⁸

This is the combination of circus movements with rolling movements familiar to those who have experimented on the brain of fish, where a destruction of one side of the midbrain calls forth rolling motions as well as circus motions toward the same side. Holmes's observations

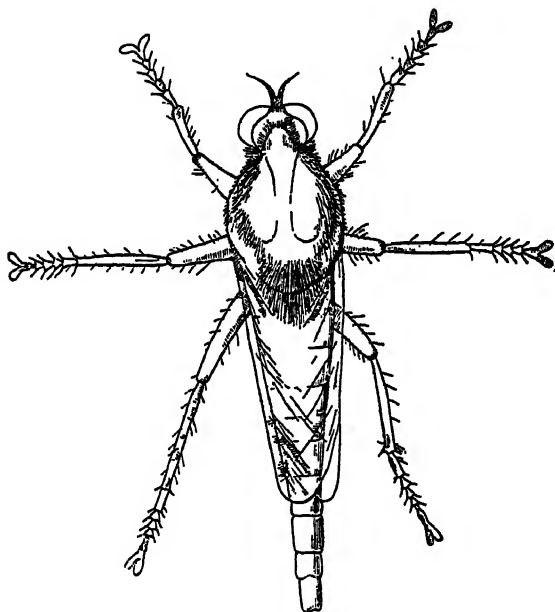


FIG. 19.—Robber fly under normal conditions seen from above. (After Garrey.)

were extended by Garrey's experiments on a large number of insects. Garrey found that the robber fly (*Proctacanthus*) (Fig. 19), which is positively heliotropic, is an unusually good object for the demonstration that the heliotropic reactions of animals are of the type of forced movements. When one eye of this fly is blackened the legs on the side of the unblackened eye are flexed and the

legs on the side of the blackened eye are more extended than normally and spread farther apart.^a The body may tilt as far toward the side of the unblackened eye as to press the legs to the table (Fig. 20). There is sometimes a

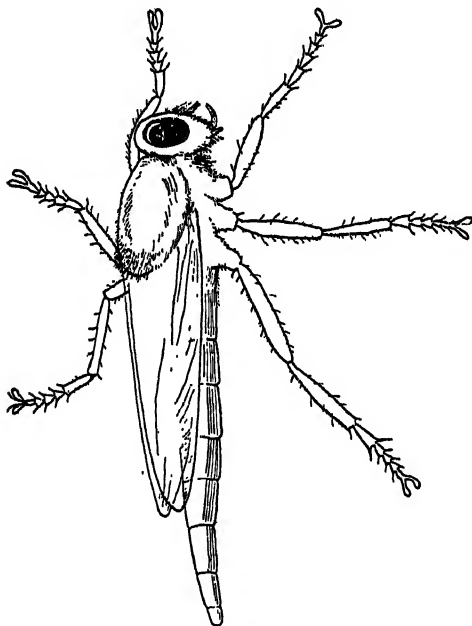


FIG. 20.—Robber fly with right eye blackened, seen from above as in Fig. 19. The body tilts over to the left side so that only the right eye is visible from above. Position of legs changed in such a way as to make motion toward left possible, toward right impossible. (After Garrey.)

tendency on the part of the body of the animal to become slightly concave toward the side of the unblackened eye.

Garrey found also that the same changes take place when one eye receives a stronger illumination than the

^a Figs. 19 to 22 and 27 were drawn from photographs kindly given to the writer for this purpose by Professor Garrey. The draughtsman was unfortunately not familiar with the anatomy of insects, which accounts for shortcomings in the drawings, which, however, have no bearing on the problem for which the drawings are intended.

other. Bringing one eye into the bright beam of light directed through the objective of the optical system of the string galvanometer, while the other eye is illuminated only by the subdued light of the optical room, promptly produced the same changes in the position of the legs and body which were observed when one eye was blackened, the more weakly illuminated eye acting like the blackened eye in the former experiment. When the illu-

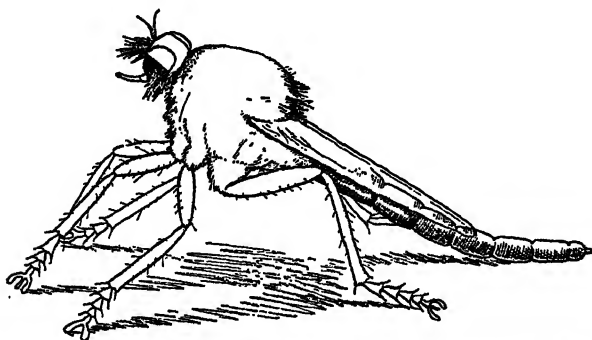


FIG. 21.—Position of robber fly when the lower halves of both eyes are blackened. Head tilted up. (After Garrey.)

mination on one side of such animals is stronger than on the other the legs on the more strongly illuminated side of the animal are bent, those on the opposite side are extended; and the head has a tendency to bend toward the light. When an impulse to move originates in the animal, it can turn easily to the light but with difficulty in the opposite direction. As soon as its head is turned to the source of light and both eyes receive the same illumination the difference in tension of the legs on the two sides of the body disappears and now the animal moves or is carried in a straight direction toward the light. By these experiments the proof of the writer's muscle tension theory of heliotropism is made complete.¹⁷⁷

Garrey observed that when the lower halves of the eyes of the robber fly are blackened the position of the legs of the two sides is symmetrical, but the anterior and middle pairs of legs are extended forward to the maximal extent, producing a striking posture in which the anterior end of the robber fly is pushed up and back from the surface of the table. The body is in opisthotonus, with the abdomen concave on the dorsal side, while the head is tilted far up and back (Fig. 21).

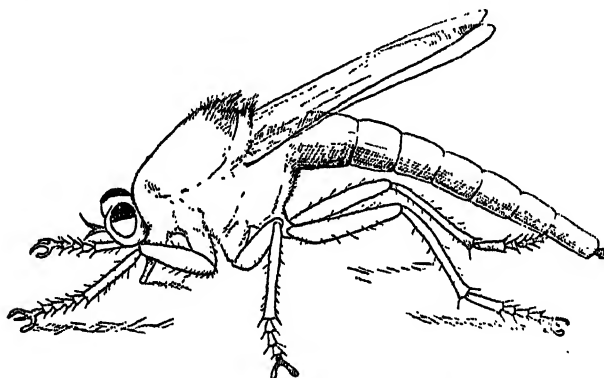


FIG. 22.—Position of robber fly when upper halves of both eyes are blackened. Head down, body convex above. (After Garrey.)

When walking these robber flies gave the impression of trying to climb up into the air. The wings are frequently somewhat spread and the animal may push itself up and back until poised vertically on the tips of the wings and abdomen. The tendency to fly is very pronounced in this condition and upon the slightest disturbance the fly soars upward and backward, striking the top of a confining glass dish or completing a circle by "looping the loop" backward. If it falls upon its back it rights itself by turning a backward somersault. Unequal blackening of the lower parts of the two eyes results in a combination of the effects just described, with those described for blackening one eye, for the animal also performs circus motions.

With the upper halves of the eyes blackened the attitude is the reverse of that described in the preceding section (Fig. 22). The anterior and middle pairs of legs are flexed. The anterior and posterior

ends of the body bend ventrally with the body in emprosthotonus. The head is bent far down. The animal may actually stand on its head, but the abdomen retains its ventral curvature, leaving a considerable angle open between its dorsum and the wings which normally rest on it.

In both walking and flying it continually keeps close to the table, and upon encountering an obstacle it frequently does a forward somersault. If it gets on its back it rights itself with greatest difficulty as its efforts simply result in bending the tail and head ventrally until they may form a complete ring. In galvanotropism the same general picture is presented by *Palæmonetes* and *Amblystoma* when the anode is at the head end, the tonus changes involved being identical in the two conditions (Garrey¹⁷⁷).

These experiments leave no doubt that the primary effect of light consists in changes in the tension of muscles and that the heliotropic reactions which appeared to the older observers as voluntary acts are in reality forced movements.

In the chapter on forced movements after brain lesion the fact was mentioned that a dog which had shown circus movements to the left after lesion of the left cerebral hemisphere shows circus motions to the right when afterward the right hemisphere is injured symmetrically; instead of being a physiologically symmetrical animal again after the second operation. The explanation is that the new operation is more effective than the old one whose effect has partly worn off. Garrey has made an observation on heliotropism which shows some analogy with this experiment on the brain.

He found¹⁷⁷ that "robber flies with one eye blackened show the postural conditions in the most pronounced way in the early morning or after being kept for some hours in the dark. Constant exposure to the light produces considerable fatigue of the eye with recovery in the dark. These facts among others suggested the possi-

bility of producing a different sensitiveness of the two eyes and corresponding differences in the muscle tonus with asymmetry of position, and in physiological action of the muscles of the two sides of the body when the two eyes were equally illuminated. Such an experiment constitutes a crucial test of the tonus theory of heliotropism. It succeeded beyond our greatest expectations. Asphalt black was applied to the right eye of several specimens of *Proctacanthus*. In two or three days the paint had formed a brittle shell. During this time the blackened eye had become 'dark adapted.' When such a fly is exposed to light, it tilts and circles to the left. If now the brittle shell is cracked off the right eye by carefully pinching with fine forceps, the exposure of this very sensitive eye to light results in a reversal of the whole picture; the fly circles toward the side from which the black was removed. Although the illumination of the two eyes is of equal intensity, what was the normal eye now becomes relatively a darkened eye owing to its lesser sensitiveness. A differential effect results, probably due to a difference in the rate of photochemical change in the two eyes. This reversal of the muscle tonus and of forced motions may persist for an hour or two or even longer, until the two eyes become, as they ultimately do, of equal sensitiveness and the fly behaves like a normal animal.

"These experiments are not only incompatible with any 'avoidance' idea, for after removal of the black there is nothing to avoid, but they are also incompatible with the conception of 'habit formation,' for 'habit' in the performance of the circling movements is of no avail when light is admitted to the darkened eye—the animals circle to that side because the tonus of the muscles is such that they are forced to do so.

“All the experiments show that the muscle tone is dependent upon the intensity of the light and that the postures assumed depend upon the relative difference in the light stimulus to the eyes. In animals with one eye completely covered the radii of the circles in which they moved were shorter the more intense the illumination of the normal eye. With one eye partially covered the circles were larger than when completely covered, and in the same way the circles were larger when one eye was covered by a film of collodion or of brown shellac, which admits some light, than when subsequently covered by opaque asphalt black. When one eye was partially covered by central application of the black paint the tilting and circling to the opposite side were abolished or reversed by brilliant illumination of the partially blackened eye. These results explain why a positively heliotropic animal with one eye blackened approaches a light by a series of alternating small and large circles, the former being executed when the good eye is illuminated from the source of light, the larger when it is in the shadow.”

We have thus far discussed chiefly positively heliotropic animals, *i.e.*, animals which are compelled to move toward the source of light. The difference between these and negatively heliotropic animals is that the legs on the illuminated side of a negatively heliotropic animal are extended, while those on the opposite side are in flexed position. This has been directly observed by Holmes, who also made sure of the fact that negatively heliotropic animals, when one eye is blackened, turn in circles with the blackened eye toward the center of the circle ²²⁸; while positively heliotropic animals turn in circles with the unblackened eye toward the center of the circle.

3. HELIOTROPISM OF UNICELLULAR ORGANISMS

In unicellular organisms, where cilia act as locomotor organs, it can easily be shown that the orientation by light is of the nature of changes in the position of cilia; this is for instance the case in respect to *Volvox*. Holmes²²⁶ states for the heliotropic reactions of this organism, that they are due to differences in the activity of the cilia on both sides of the organism and this explanation agrees with the actual observations of Bancroft on the galvanotropic reactions of *Volvox*.

In flagellates, the mechanism of locomotion is very complicated and does not consist in an oar-like action of a flagellum as was formerly assumed. Bancroft has shown that in *Euglena*, as already stated, the flagellum inserted at the anterior end of the organism is bent backward in the form of an inverted U, and that locomotion is brought

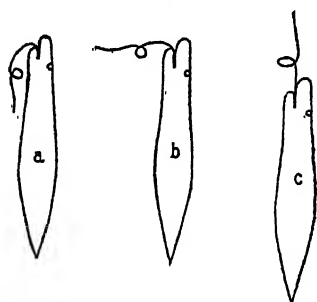


FIG. 23.—Diagram showing the position of the flagellum as seen in a viscid medium. *a*, when *Euglena* is swimming forward in a narrow spiral; *b*, when swerving sharply toward the dorsal side; *c*, when moving backward. (After Bancroft.)

about by the formation of a loop which travels from the base of the flagellum toward the free end (Fig. 23). The path of the organism which results from this action is a spiral with continual rotation of the organism around its longitudinal axis. Bancroft has shown that the behavior of the organism under the influence of light is identical with that in a constant galvanic field.²¹ One-sided illu-

mination as well as a current going transversally through such an organism cause changes in the position of cilia comparable with those observed in the legs of crustaceans, insects, and vertebrates.

4. HELIOTROPISM OF SESSILE ANIMALS

When we study the effects of light on sessile animals we find that they behave in a similar manner to sessile plants. When illuminated from one side they bend their heads to the source of light until their axis of symmetry goes through the source of light. In this case the symmetrical photosensitive elements receive equal illumination and the symmetrical muscles are under equal tension. Hence the animal remains in this orientation. These sessile animals were the first examples by which the



FIG. 24.—Tube worms in aquarium, all bending toward light.

muscle tension theory of animal heliotropism was proved.²⁸⁸

Spirographis spallanzani (Fig. 24) is a marine annelid from 10 cm. to 20 cm. long, which lives in a rather rigid yet flexible tube. The latter is formed by a secretion from glands at the surface of the animal. The tube is attached by the animal with its lower end to some solid body, while the other end projects into the water. The worm lives in the tube and only the gills, which are arranged in a spiral at the head end of the worm, project from the tube. The gills, however, are quickly retracted, and the worm withdraws into the tube when touched or if a shadow is cast upon it.

When such tubes with their inhabitants are put into an aquarium which receives light from one side only, it requires, as a rule, a day or more until the foot end of the tube is again attached to the bottom of the aquarium. As soon as this occurs, the anterior end of the tube is raised by the worm until the axis of symmetry of the gills falls into the direction of the rays of light (Fig. 24) which enter through the window into the aquarium.²⁸⁸ When the animal has once reached this position it retains it as

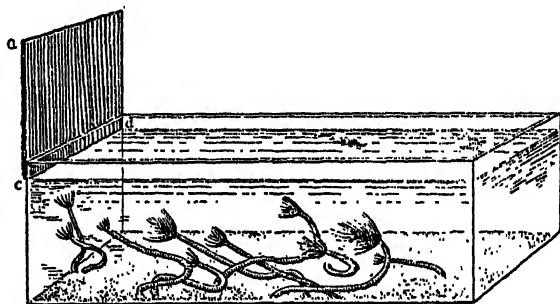


FIG. 25.—The same animals after the position of the aquarium to the window was reversed.

long as the position of the aquarium and the direction of the rays of light remain the same. When, however, the aquarium is turned 180° , so that the light falls in from the opposite direction, the animal bends its tube during the next twenty-four or forty-eight hours in such a way that the axis of symmetry of its circle of gills is again in the direction of the rays of light (Fig. 25). When the light strikes the aquarium from above, the animals assume an erect position, like the positively heliotropic stems of plants when they grow in the open.

In these phenomena the mechanical properties of the tube play a rôle. When the animal is taken out of the bent tube, the latter retains its form. This permanent

change of form of the tube is apparently caused through the secretion of new layers on the inside of the tube. The youngest layers of the secretion are more elastic than the old layers, and, moreover, have at first a powerful tendency to shorten. If such a secretion occurs on one side of the tube only, or more so than on the opposite side, the former must become shorter than the latter, and the result must be a curvature of the tube, that side becoming concave where the new secretion has occurred.

On this assumption the process of heliotropic curvature is in this case as follows: when the light strikes the circle of gills from one side only, in these elements certain photochemical reactions occur to a larger extent, than on the opposite side. This results in corresponding alterations of the sensory nerve endings, the sensory nerves and the corresponding motor nerves, and their muscles. The sense of these changes is such as to throw the muscles connected with the nerves of the gills on the light side into a more powerful tonic or static contraction than the muscles on the opposite side of the body. The consequence is a bending of the circle of tentacles, or the head, toward the source of light, which will continue until the axis of symmetry of the circle of tentacles falls into the direction of the rays of light. When this happens, symmetrical tentacles are struck at the same angle (or in other words, with equal intensity) by the rays of light, and therefore the tone of the antagonistic muscles is the same. The result is that the circle of tentacles becomes fixed in this position. The bending of the head produces an increased pressure and friction of the animal against that side of the tube which is directed toward the light, and this pressure and friction lead to an increased secretion and the formation of a new layer inside the tube.

Heliotropic curvature of sessile animals can be shown equally well in a hydroid, *Eudendrium*. It is necessary to cut off the old polyps at once when the animal is brought into the laboratory and to put the stem into fresh, clear,

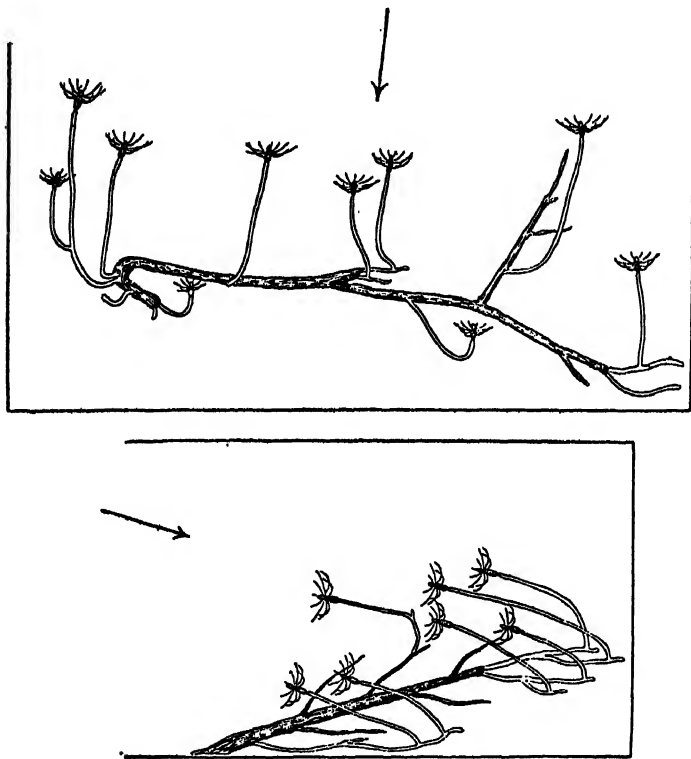


FIG. 26.—Polyps of *Eudendrium*, all growing toward source of light. The arrow indicates the direction of the rays of light, which in one case fall in from above, in the other from the left side.

sea water. In a day or two new polyps are formed by regeneration and these polyps will bend toward the light until their axis of symmetry is in the direction of the rays of light (Fig. 26). The region at the base of the polyps is contractile and when light strikes the polyps from one

side only, the stem on the side of the light contracts more than on the other side, and this results in a bending of the stem, whereby the polyp is put into the direction of the rays of light. As soon as the axis of the polyp is in the direction of the rays of light (provided there is only one source of light), the tension of the contractile elements is the same all around, and there is no more reason for the organism to change its orientation. It, therefore, remains in this orientation to the light.

The muscle tension theory of animal heliotropism is, therefore, proved for all classes of the animal kingdom, infusorians, hydroids, annelids, crustaceans, etc. It would be wrong to state that the theory holds only for insects.

CHAPTER VI

AN ARTIFICIAL HELIOTROPIC MACHINE

THE reader will have perceived that in the preceding analysis animals are treated as machines whose apparently volitional or instinctive acts, as *e.g.*, the motion toward the light, are purely physical phenomena. The best proof of the correctness of our view would consist in the fact that machines could be built showing the same type of volition or instinct as an animal going to the light. This proof has been furnished by the well-known inventor, Mr. John Hays Hammond, Jr. The following is a description of the machine given by one of Mr. Hammond's fellow-workers who coöperated with him in the development of the machine, Mr. B. F. Miessner.

This "Orientation Mechanism" consists of a rectangular box, about 3 feet long, 1½ feet wide, and 1 foot high. This box contains all the instruments and mechanism, and is mounted on three wheels, two of which are geared to a driving motor, and the third, on the rear end, is so mounted that its bearings can be turned by solenoid electro-magnets in a horizontal plane. Two 5-inch condensing lenses on the forward end appear very much like large eyes.

If a portable electric light, such as a hand flashlight, be turned on in front of the machine it will immediately begin to move toward the light and, moreover, will follow that light all around the room in many complex manœuvres at a speed of about 3 feet per second. The smallest circle in which it will turn is about 10 feet diameter; this is due to the limiting motion of the steering wheel.

Upon shading or switching off the light the "dog" can be stopped immediately, but it will resume its course behind the moving light so long as the light reaches the condensing lenses in *sufficient intensity*. Indeed, it is more faithful in this respect than the proverbial ass behind the bucket of oats. To the uninitiated the performance of the pseudo dog is very uncanny indeed.

The explanation is very similar to that given by Jacques Loeb, of reasons responsible for the flight of moths into a flame. . . .

The *orientation mechanism* here mentioned possesses two selenium cells corresponding to the two eyes of the moth, which when influenced by light effect the control of sensitive relays instead of controlling nervous apparatus, as is done in the moth. The two relays (500 to 1,000 ohm polarized preferred) controlled by the selenium cells in turn control electro-magnetic switches, which effect the following operations: When one cell or both are illuminated the current is switched on to the driving motor; when one cell alone is illuminated an electro-magnet is energized and effects the turning of the rear steering wheel. The resultant turning of the machine will be such as to bring the shaded cell into the light. As soon and as long as both cells are equally illuminated in sufficient intensity, the machine moves in a straight line toward the light source. By throwing a switch, which reverses the driving motors, the machine can be made to back away from the light in a most surprising manner. When the intensity of the illumination is so decreased by the increasing distance from the light source that the resistance of the cells approach their dark resistances, the sensitive relays break their respective circuits and the machine stops.

The principle of this orientation mechanism has been applied to the "Hammond Dirigible Torpedo" for demonstrating what is known as *attraction by interference*. That is, if the enemy tries to interfere with the guiding station's control the torpedo will be attracted to him, etc.^a

Nothing seems to have been published beyond these meagre details, but the writer understands that the active machine has been demonstrated in a number of places in this country. It seems to the writer that the actual construction of a heliotropic machine not only supports the mechanistic conception of the volitional and instinctive actions of animals but also the writer's theory of heliotropism, since this theory served as the basis in the construction of the machine. We may feel safe in stating that there is no more reason to ascribe the heliotropic reactions of lower animals to any form of sensation, *e.g.*, of brightness or color or pleasure or curiosity, than it is to ascribe the heliotropic reactions of Mr. Hammond's machine to such sensations.

^a *Electrical Experimenter*, September, 1915, 202.

CHAPTER VII

ASYMMETRICAL ANIMALS

It was necessary for us to begin our analysis with symmetrical animals since as the result of this analysis the conduct of asymmetrical organisms offers no difficulty. The result of the asymmetry consists merely in a change in the geometrical character of the path in which an animal is compelled to move to or from the source of energy. While this path is a straight line in a symmetrical and positively heliotropic organism it is a spiral around this straight line as an axis in an asymmetrical organism, like *Euglena*. Suppose a positively heliotropic animal to have slightly asymmetrical appendages which give it a tendency to deviate to the left. Let us suppose that the plane of symmetry of the animal goes at the beginning of the experiment through the source of light and that the animal is swimming toward the light. After a few strokes the head of the organism will have deviated slightly to the left on account of the asymmetry in the activity of the appendages. As soon as the median plane of the animal deviates to the left, the left eye is less illuminated than the right one. As a consequence, a difference in the tension of the muscles on the two sides of the animal will be produced which will compensate the natural lack of symmetry in the muscles and the animal will cease to deviate further to the left; and this compensating effect of the unequal illumination of the two eyes will continue until the animal is actually oriented in the right way again, *i.e.*,

until its plane of symmetry goes through the source of light. All that the inherited or accidental asymmetry does is to cause the animal to move in a path which is not a mathematically straight line; but this deviation will be marked only in a case of very pronounced or excessive asymmetry.

We have already described the behavior of a dog whose left cerebral hemisphere has been injured and who has a tendency to deviate to the left. When such a dog is shown a piece of meat it moves toward it in a fairly straight line, its tendency to deviate to the left being compensated by the orienting effect of the retina image of the piece of meat. If the dog deviates to the left, the piece of meat is apparently dislocated to the right of the dog and this dislocation alters the tension of the muscles on the two sides of the animal in such a way as to make it turn back to the right. In this way the dog reaches the piece of meat in a fairly straight line, though with a greater amount of labor, since the tendency to deviate to the left is constantly compensated automatically by a stronger contraction of the muscles turning the animal to the right.

The writer showed many years ago that many insects have a tendency to creep upward, and that this is due to an orienting effect of gravity upon the animal. When a perfectly symmetrical insect is put on a vertical stick it walks upward in a straight line. What will happen when such an animal is made asymmetrical? Garrey has performed this experiment by using flies in which one eye was blackened. As we have seen, such organisms are rendered asymmetrical not only in regard to the eyes but also in regard to their apparatus of locomotion, since in one side of the body the tension of the flexors, in the

legs of the other side the tension of the extensors prevails. As a consequence the fly has a tendency to move in circles with the intact eye toward the center.

Garrey has shown that when a fly with one eye blackened is put on a vertical stick, it still walks upward, but in



FIG. 27.—Fly with one (right) eye blackened can creep only in a spiral on a vertical stick, while normally it creeps in a straight line. (After Garrey.)

spirals around the stick (Fig. 27), instead of in a straight line. The asymmetry of locomotion changes only the geometrical nature of the path in which the animal moves, from a straight line to a spiral, but does not alter the forced movement character of the reaction.

Bancroft has pointed out that when in a positively heliotropic amphipod one eye is blackened and the legs of the same side are cut off, the animal's path would be a combination of a circus motion induced by the blackening of the eye and of a rolling motion around its longitudinal axis. Both effects combined would result in the animal swimming in a spiral path, and if the animal is positively heliotropic it would swim in such a path toward the

light. This is the path which aquatic, asymmetrical positively heliotropic organisms, such as the flagellate *Euglena*, describe in their motions to the light.

But this locomotor mechanism (of *Euglena*) is imperfect, it forces the organism to move in a spiral, and always to turn toward a structurally determined side. There are many organisms which swim in spirals and become oriented by turning toward a structurally defined side. Jennings and Mast include all such orientations under "trial and error" and contrast them with the direct orientation of such animals as the amphipods in which the turning may be either toward the left or the right. Let us now consider whether the orientation of *Euglena* is more like the selection of random movements (which we would all agree may justifiably be called "trial and error"), or whether it is more like the orientation of the terrestrial amphipods studied by Holmes.

I think that all students of behavior including Jennings and Mast believe that in the case of these amphipods we have direct heliotropic orientation. If the right eye of such a positively heliotropic amphipod be covered with asphalt varnish it will execute circus movements towards the left. The usual explanation is that the main nervous connection is between the eye on one side and the legs on the opposite side of the body. The light shining on the uncovered eye brings about a condition of increased muscular tonus in the legs of the opposite side, which is not present in the legs connected with the covered eye. Consequently the right legs push more strongly and the amphipod turns towards the left.

Suppose now we remove some or all of the left legs from an amphipod of this kind so that it will always turn toward the left, and transfer it to water in which it must be supposed to swim in a spiral path. We will then have an organism which would become oriented in essentially the same way that *Euglena* does. The animal would always swerve toward the left. But, when the spiral course brings it into such a position that the light shines directly on the left eye, the muscular tonus of the right legs would be increased and the swerving toward the light would increase. Thus orientation would be effected in just the same way that it is in *Euglena*.

While these hypothetical changes that must be made in the amphipod, to make it react like *Euglena*, are considerable, they concern only the details. The fundamental nature of the photochemical substances, the nature of their stimulation and the character of their connection with the locomotor organs have none of them been modified. All that has been done is to make an asymmetrical organism swimming in a spiral out of a bilateral one.^a These changes are much less fundamental than

^a Swimming in a straight line.

those which we would have to imagine in order to make an amphipod orient to light by the selection of random movements. In order to bring about this latter change the whole nature of the photochemical substances and their relations to the leg muscles would have to be modified. In the one case the required changes are all of a mechanical nature and so simple that the experiment might possibly succeed. In the other case the required changes are largely chemical, and so complex that we have no data for even imagining what ought to be done in order to bring them about (Baneroff ²¹).

The asymmetry of organisms only modifies the geometrical character of the path but not the mechanism of the reaction.

CHAPTER VIII

TWO SOURCES OF LIGHT OF DIFFERENT INTENSITY

THE writer observed that if heliotropic animals are exposed to two equidistant lights of equal intensity they move in a line perpendicular to the line connecting the two lights.^{287, 294} This has been confirmed by numerous observers, *e.g.*, Bohn on *Littorina*, by Parker and his pupils, especially by Bradley M. Patten on the larvæ of the blowfly, by Loeb and Northrop on the motions of the larvæ of *Balanus*, and by others. The question arises: In which line will an animal move when the intensity of the two lights differs? When the animal is positively heliotropic it should cease to move in a line at right angles to the line connecting the two lights but should move in a line which deviates toward the stronger of the two lights; if the animal is negatively heliotropic it should deviate toward the weaker of the two lights. When the two eyes are illuminated by two lights of different intensity, the illumination in both eyes can become approximately equal only when the eye struck by the weaker light is exposed at a larger angle than the eye struck by the stronger light. Under such conditions, the animal should be compelled to move in a straight line which, however, is no longer at right angles to the line connecting the two lights, but which deviates to an extent determined by the difference in the intensity of the two lights. The case was

worked out quantitatively by Bradley M. Patten on a negatively heliotropic animal, the full grown larva of the blowfly.^{412, 413} The source of light was at G (Fig. 28) (one or more Nernst lamps of measured candle power), a portion of light from these lamps passed through the screens d and d' to the mirrors M and M' , set at a definite

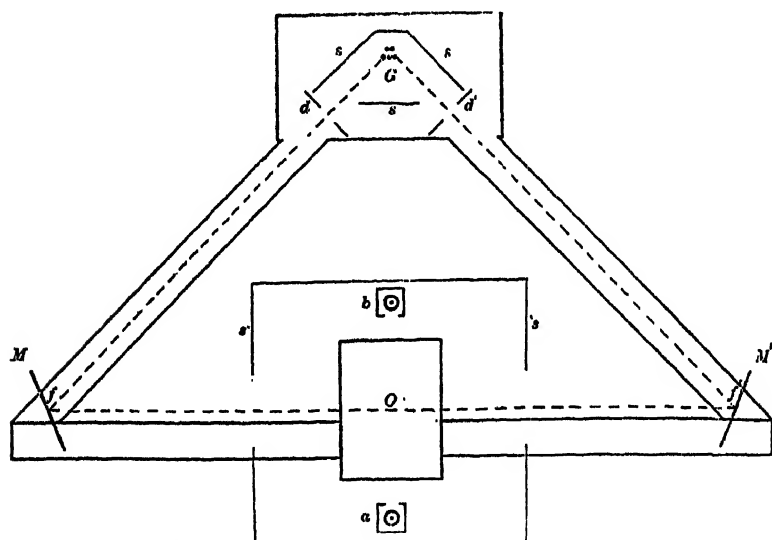


FIG. 28.—Diagram of apparatus used to produce differential bilateral light stimulation. G , five 220-volt Nernst glowers; M and M' , mirrors; f and f' , central point of mirrors; O , center of observation stage; dotted lines, central ray of beam of light from the glowers reflected to O by the mirrors; d and d' , screens with rectangular openings; s and s' , light shields; a and b , 2 c.p. orienting light with screens. (After Patten.)

angle so that the rays were reflected to the observation point O . The two beams of light reaching O were of the same intensity. With the means of one of the lights at a or b the animal was first caused to move across the field O at right angles to the rays reflected from the mirrors M and M' . The animals first started in this direction, then came suddenly under the influence of the light re-

flected by M and M' . In order to make the ratio of intensities of light from M and M' different, the observation stage O was put at unequal distance from M and M' . The larvæ were made to record their trails while moving under the influence of two lights and the deviation of this trail from the perpendicular upon the line connecting the two sources of light M and M' was measured with a goniometer (Fig. 29). The result of the measurements of 2,500 trails, showing the progressive increase in angular deviation of the larvæ (from the perpendicular upon the line connecting the two lights) with increasing differences between the lights, are given in Table I. Since the deviation or angular deflection was toward the weaker of the two lights (the animal being negatively heliotropic) the deviation is marked negative.

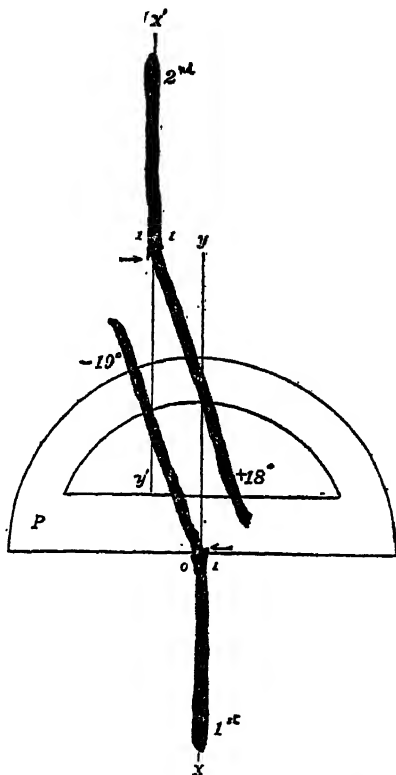


FIG. 29.—Diagram to show the method of measuring trails. The lines xy and $x'y'$ are drawn through the trails at the points reached—marked by the arrows—when the side lights were turned on. The angle of deflection from this line is measured by a protractor, P . The small figures near the arrows indicate the number of wig-wag movements made when the side lights were turned on; 1st and 2nd refer to the sequence in which the trails were run. (After Patten.)

TABLE I.

| Percentage difference in the intensity of the two lights | Average angular deflection of the two paths of the larvæ toward the weaker light |
|--|--|
| Per cent. | Degrees |
| 0 | - 0.09 |
| 8 $\frac{1}{3}$ | - 2.77 |
| 16 $\frac{2}{3}$ | - 5.75 |
| 25 | - 8.86 |
| 33 $\frac{1}{3}$ | -11.92 |
| 50 | -20.28 |
| 66 $\frac{2}{3}$ | -30.90 |
| 83 $\frac{1}{3}$ | -46.81 |
| 100 | -77.56 |

Patten also investigated the question whether the same difference of percentage between two lights would give the same deviation, regardless of the absolute intensities of the lights used (Weber's law). The absolute intensity was varied by using in turn from one to five glowers. The relative intensity between the two lights varied in succession by 0, 8 1/3, 16 2/3, 25, 33 1/3, and 50 per cent. Yet the angular deflections were within the limits of error identical for each relative difference of intensity of the two lights, no matter whether 1, 2, 3, 4, or 5 glowers were used. Table II gives the results.

TABLE II

A TABLE BASED ON THE MEASUREMENTS OF 2,700 TRAILS SHOWING THE ANGULAR DEFLECTIONS AT FIVE DIFFERENT ABSOLUTE INTENSITIES

| Number of glowers | Difference of intensity between the two lights | | | | | |
|-------------------|--|---------------------------|----------------------------|--------------|----------------------------|--------------|
| | 0 per cent. | 8 $\frac{1}{3}$ per cent. | 16 $\frac{2}{3}$ per cent. | 25 per cent. | 33 $\frac{1}{3}$ per cent. | 50 per cent. |
| | Deflection in degrees | | | | | |
| 1 | -0.55 | -2.32 | -5.27 | -9.04 | -11.86 | -19.46 |
| 2 | -0.10 | -3.05 | -6.12 | -8.55 | -11.92 | -22.28 |
| 3 | +0.45 | -2.60 | -5.65 | -8.73 | -13.15 | -20.52 |
| 4 | -0.025 | -2.98 | -6.60 | -9.66 | -11.76 | -19.88 |
| 5 | -0.225 | -2.92 | -5.125 | -8.30 | -10.92 | -19.28 |
| Average... | -0.09 | -2.77 | -5.75 | -8.86 | -11.92 | -20.28 |

On the writer's theory the following explanation of these deviations should be given. The muscles moving the head of the animal to the side of the weaker illumination, having a higher tension than their antagonists, bring about a deflection of the animal toward the side of the weaker light. As soon as its two photosensitive areas in the head—the animal has no eyes—which are not parallel, but inclined to each other are deflected from the perpendicular upon the line connecting the two lights, the photosensitive areas of the animal will no longer be struck by the lights at the same angle, but on the side of the weaker light the area will be struck at an angle nearer to 90° than the photosensitive area exposed to the stronger light. In this way the change in angle will compensate the difference in intensity of the two lights until the orientation of the animal is such that the compensation is complete and both photosensitive areas receive the same illumination. The animal will then continue to move in this direction.

Patten has computed the angle of the photosensitive surfaces for these animals from the angle of their orientation under varying inequalities of illumination.

This angle has been computed for the blowfly larva, using the "angular deflections" already ascertained. The magnitude of the angle may bear no direct relation to the actual angle at which the sensitive areas are located in the body of the animal, because of the many factors which may modify the direction of the rays before they fall on the sensitive surfaces. The significant test of the hypothesis would be the constancy of the angle when computed from experimental data obtained under varying conditions.

The method of constructing such an angle is shown in Fig. 30, in which the opposing lights are assumed to be of a two-to-one ratio of intensity. The line *AB* is drawn perpendicular to the direction of the rays of light. On the line *AB*, construct angle *BOC* equal to the actual average angular deflection of the larvæ at a two-to-one ratio of lights.

to AB . Lay off on AB distances hx and hy , such that $hy = 2hx$. From x and y erect lines perpendicular to AB , they will intersect OC at f and e respectively. Bisect the line ef , and at its middle point, g , construct a line kl perpendicular to OC . From the point of intersection of kl and yy' (M), draw a line to D . From the intersection of kl and xx' (N), draw a line to D .

The angle MDN is the desired angle.

Proof: $eg = gf$ (construction).

Angle $egM = \text{angle } fgN$ (construction).

Angle $Meg = \text{angle } Nfg$ (alternate int. angles of parallel lines, yy' and xx' being parallel by construction).

Therefore triangle $Meg = \text{triangle } Nfg$ (side and two adjacent angles being equal).

$Ng = gM$ (similar sides of equal triangles).

$gD = gD$ (identical).

Therefore triangle $NgD = \text{triangle } MgD$ (rt. triangles, altitude and base equal).

Therefore angle $gDM = \text{angle } gDN$ and side $DM = \text{side } DN$.

Now by construction hx is the projection of DN on AB and hy the projection of MD on AB , and by construction $hy = 2hx$.

This fulfills all the conditions of construction.

The equal lines MD and DN represent equal bilateral sensitive areas inclined to each other at such an angle, MDN , that the surface represented by MD intercepts an area of light twice as great as the surface represented by DN , its projection on the perpendicular to the light rays being twice as great ($hy = 2hx$). But the light falling on DN is of twice the intensity of the light falling on DM , so that the total amount of light received by each of the equal areas is the same.

By this method of construction, the average angle of sensitiveness was computed for four intensity differences, using as a basis the angular deflection of the larvæ as determined by experiment. The magnitude of the angles is almost identical in all four cases.⁴¹²

Experiments by a somewhat different method, to be discussed in the next chapter, on the positively heliotropic larvæ of the barnacle show that these results of Patten are more general.

We may, therefore, say that the migration of animals to or from the light is of the nature of a forced movement determined by the effect of light on the photosensitive elements of the body. Unequal illumination of symmetri-

cal photosensitive elements on the two sides of the body alters the tension of symmetrical muscles, and as a consequence the animal is, when moving, compelled to change its direction of motion until it is oriented in such a way to the light that symmetrical elements receive the same illumination. In this case the tension of symmetrical muscles is equal again and the animal is compelled to move in this direction.

It has been suggested by the anthropomorphic interpreters of animal conduct that the motion of an animal to a source of light is the same phenomenon as when a human being who has lost his way in the dark is attracted by an illuminated human habitation. As Bohn pointed out, the definite path in which a positively heliotropic animal moves when under the influence of two lights, shows that the anthropomorphic interpretation is as erroneous in this as in any other case. A human being would go to one of two illuminated houses and not toward a point between them, determined by the relative intensity of the two lights.⁶⁸

CHAPTER IX

THE VALIDITY OF THE BUNSEN-ROSCOE LAW FOR THE HELIOTROPIC REACTIONS OF ANIMALS AND PLANTS

WE have thus far said little about the identity of the heliotropism of plants and animals. Yet the two phenomena are essentially alike. When we keep positively heliotropic sessile plants and sessile animals near a window, both will bend toward the source of light, though the mechanism of bending may not be the same in all details, the bending being produced in the case of the plant (and possibly in certain animals like *Eudendrium*) by unequal growth in length of the plant on the illuminated and shaded sides; while in the case of higher animals, e.g., *Spirographis*, it is produced by differences in the tension of the muscles on the illuminated and shaded sides of the animal. Motile plant organisms like *Volvox*, are driven to the source of light, owing to differences in the tension of the contractile organs on the shaded and illuminated side, and the same is true for animals like insects.

A further point of coincidence lies in the validity of the photochemical law of Bunsen and Roscoe for the heliotropism of animals and plants.

The law of Bunsen and Roscoe says that within certain limits the chemical effect produced by light increases in proportion with the product of intensity into the duration of illumination, e.g., $Effect = Kit$, where i is intensity, t duration of illumination, and K a constant. This is true

for the blackening of photographic paper by light, and it can be shown that the same law holds for heliotropic reactions of plants as well as animals.

Blaauw^{46,47} established this fact for the etiolated seedlings of *Avena sativa*. These organisms were exposed to lights of a definite candle power for some time and then left in the dark. After a certain time the seedlings began to bend, becoming concave on that side which had previously been illuminated. By varying the candle power of light (*i*) and the duration of illumination (*t*), he found that the value of *it* required to cause 50 per cent. of the seedlings to bend was always the same. Table III gives

TABLE III

Time required for different intensities of light to produce heliotropic curvatures in 50 per cent. of the seedlings of *Avena*

| Candle-meter | Duration of illumination | Candle-meter-seconds |
|--------------|--------------------------|----------------------|
| 0.00017 | 43 hours | 26.3 |
| 0.000439 | 13 hours | 20.6 |
| 0.000609 | 10 hours | 21.9 |
| 0.000855 | 6 hours | 18.6 |
| 0.001769 | 3 hours | 19.1 |
| 0.002706 | 100 minutes | 16.2 |
| 0.004773 | 60 minutes | 17.2 |
| 0.01018 | 30 minutes | 18.3 |
| 0.01640 | 20 minutes | 19.7 |
| 0.0249 | 15 minutes | 22.4 |
| 0.0498 | 8 minutes | 23.9 |
| 0.0898 | 4 minutes | 21.6 |
| 0.6156 | 40 seconds | 24.8 |
| 1.0998 | 25 seconds | 27.5 |
| 3.02813 | 8 seconds | 24.2 |
| 5.456 | 4 seconds | 21.8 |
| 8.453 | 2 seconds | 16.9 |
| 18.94 | 1 second | 18.9 |
| 45.05 | 2/5 seconds | 18.0 |
| 308.7 | 2/25 seconds | 24.7 |
| 511.4 | 1/25 seconds | 20.5 |
| 1,255 | 1/55 seconds | 22.8 |
| 1,902 | 1/100 seconds | 19.0 |
| 7,905 | 1/400 seconds | 19.8 |
| 13,094 | 1/800 seconds | 16.4 |
| 26,520 | 1/1000 seconds | 26.5 |

the time required for different intensities of light varying from 0.00017 to 26,520 candle power to cause 50 per cent. of the seedlings to show heliotropic curvatures. As can be seen, the product *it* is always approximately 20.

Ewald and the writer^{300, 305} tested the validity of the law of Bunsen and Roscoe for the heliotropic curvatures of *Eudendrium*. A number of stems of *Eudendrium*, from which the polyps had been cut off, were put upright into a trough with parallel walls, containing sea water. As soon as the new polyps had regenerated they were exposed to light of a certain intensity for a short time and then kept in the dark. In the dark the bending of the polyps in the direction of the former source of light occurred. The purpose was to find the minimum time of exposure required for a given light (40 candle power) to induce 50 per cent. of the polyps to bend to the light (Table IV).

TABLE IV

| Percentage of polyps bending toward the former source of light | | | | | |
|--|---|------|--------|------|------|
| Duration of illumination | Distance of the polyps from the light in meters | | | | |
| | 0.25 | 0.50 | 1.00 | 1.50 | 2.00 |
| 10 | 65 ¹ | | | | |
| 15 | 68 | | | | |
| 20 | 74 | | | | |
| 30 | | 42 | | | |
| 35 | | | | | |
| 40 | | 56 | | | |
| 45 | | 60 | | | |
| 50 | | | | | |
| 60 | | 60 | | | |
| 90 | | | | | |
| 120 | | 65 | 30 | | |
| 150 | | | 48, 50 | | |
| 180 | | | | | |
| 240 | | | | | |
| 300 | | | 85 | 40 | |
| 360 | | | | 40 | |
| 420 | | | | 57 | (15) |

¹ Very young, abnormally sensitive polyps.

If we calculate from this the value of the product *it* for different intensities of light we find that it obeys the Bunsen-Roscoe law (Table V).

TABLE V

| Distance of polyps from light | Time required to call forth heliotropic curvature in 50 per cent. of the polyps | |
|-------------------------------|---|---|
| | Observed | Calculated according to the Bunsen-Roscoe law |
| <i>Meters</i> | <i>Minutes</i> | <i>Minutes</i> |
| 0.25 | 10 | |
| 0.50 | between 35 and 40 | 40 |
| 1.00 | 180 | 160 |
| 1.50 | between 360 and 420 | 360 |

The material varies considerably so that it is not always possible to induce 50 per cent. of the polyps to undergo heliotropic curvature. For this reason Loeb and Wasteneys³¹² repeated these experiments by a somewhat different method.

We confined our experiments to three intensities of light by putting the specimens at distances of 25, 37.5, and 50 cm. from a Mazda incandescent lamp, of about 33 Hefner candles. The times of exposure were adjusted so that on the assumption of the applicability of the Bunsen-Roscoe law the same effect, *i.e.*, the same percentage of polyps bending towards the light should be produced. Thus in some experiments the exposure for the three distances given was 10, 22.5, and 40 minutes respectively, in others, 7, 15.75, and 28 minutes, and so on. The ratios of the percentage of polyps bending toward the light for the three distances should be as 1:1:1. Since the material differed widely in different experiments and in different dishes, it was necessary to compute the averages of a large number of experiments.

The colonies, immersed in sea water, were arranged

in a row in rectangular glass dishes, the stems being inserted in holes made in a layer of paraffin mixed with lamp black as in the previous experiments. The rear side of the dish was also coated with the paraffin lamp black mixture in order to prevent reflection of light from the slightly uneven back surface of the dish.

Table VI gives a summary of the results. The first three columns give the times of exposure for the three

TABLE VI

| Times of exposure in minutes | | | Ratio of per cent. of hydranths bending towards light | | |
|------------------------------|----------|--------|---|----------------|------------------|
| 25 cm. | 37.5 cm. | 50 cm. | 25 cm.: 37.5 cm. | 25 cm.: 50 cm. | 37.5 cm.: 50 cm. |
| 15 | | 60 | | 1.50 | |
| 20 | | 80 | | 1.30 | |
| 10 | 22.5 | 40 | 1.20 | (3.08) | (2.56) |
| 10 | 22.5 | 40 | 0.94 | 1.47 | 1.55 |
| 10 | 22.5 | 40 | 1.57 | (2.30) | (2.43) |
| 10 | 22.5 | 40 | 1.43 | 1.04 | 0.94 |
| 10 | 22.5 | 40 | 0.76 | 1.09 | 1.47 |
| 10 | 22.5 | 40 | 1.05 | 1.13 | 0.90 |
| | | | | | 0.96 |
| 10 | 22.5 | 40 | 1.15 | | 0.99 |
| 7 | 15.75 | 28 | 0.84 | 0.62 | 0.74 |
| 7 | 15.75 | 28 | 1.70 | 0.49 | 0.58 |
| 7 | 15.75 | 28 | 0.85 | 1.25 | 1.35 |
| 7 | 15.75 | 28 | (2.09) ¹ | 0.99 | 1.08 |
| 7 | 15.75 | 28 | 1.14 | 1.15 | 0.55 |
| 7 | 15.75 | 28 | 0.44 | 0.92 | 0.44 |
| 7 | 15.75 | 28 | 1.52 | 0.80 | 0.61 |
| 7 | 15.75 | 28 | 0.59 | 0.36 | 0.70 |
| 7 | 15.75 | 28 | 0.48 | 1.07 | 0.31 |
| 7 | 15.75 | 28 | 1.00 | 0.48 | 1.80 |
| 7 | 15.75 | 28 | 0.69 | 1.09 | 0.81 |
| 7 | 15.75 | 28 | 1.26 | 0.85 | 1.09 |
| 7 | 15.75 | 28 | 0.86 | 1.38 | 0.85 |
| 7 | 15.75 | 28 | 0.70 | 1.07 | 1.59 |
| 7 | 15.75 | 28 | | 0.77 | 1.24 |
| 7 | 15.75 | 28 | | 0.60 | |
| Mean..... | | | 1.02 | 0.99 | 1.02 |
| Probable error.. | | | ±0.01 | ±0.01 | ±0.01 |

¹ Bracketed values being extreme variates are excluded from calculations of the means and probable errors.

distances of the source of light, selected, as stated, on the assumption that the Bunsen-Roscoe law holds. On that assumption the ratio of percentage bent in any two or all three dishes on any one day should equal 1.0. These ratios for each pair of distances of the source of light are given in the three other columns of the table. The percentage bending was only compared in dishes containing material regenerated and exposed on any one day, since only in this case was there any likelihood that the material was in any way uniform, and since only in this case the experiments were carried on at the same temperature and the same conditions of regeneration.

The result was that the observed ratios were as 1.02:0.99:1.02 (with a probable error of ± 0.01) while the values calculated on the assumption of the validity of the Bunsen-Roscoe law were as 1:1:1; *i.e.*, the results showed as great an approximation between observed and calculated values as one could expect.

There is a second method for testing the validity of the Bunsen-Roscoe law, based on the use of two sources of light of equal intensity.

If it is true that the heliotropic efficiency of light is determined by the product of intensity, i , into duration of illumination, t , we can alter this product by varying t as well as by varying i .

Rádl had shown that the position of the eye of the fresh water crustacean, *Daphnia*, is determined by the position of a source of light,⁴⁴⁷ and Ewald¹⁴⁵ found that by exposing the eye to two different sources of light simultaneously the eye is put into a position determined by the relative intensity of the two lights. When one light remained constant and the intensity of the other light was lowered the position of the eye changed. He now

could show that when the duration of illumination of one eye was altered by a rotating opaque disk with one sector cut out, the heliotropic effect on the eye of *Daphnia* was the same as when the intensity i of the same light was reduced to an amount corresponding to the Bunsen-Roscoe law.

Under the influence of two constant lights of equal intensity heliotropic animals move in a direction at right angles to the line connecting the two lights. If the law of Bunsen and Roscoe holds the effect of a constant light should be diminished if a rapidly rotating opaque disk with one sector cut out be put in front of the light, and the diminution should be equal to the fraction of the arc of the sector. Thus a sector of 90° , which reduces the total duration of illumination to one-fourth, should also reduce the heliotropic effect of the light to one-fourth, and the animal should deviate from the old direction in the direction toward the light without a disk before it. If, however, we lower the intensity of the latter light to one-fourth by doubling its distance we also reduce its heliotropic effect to one-fourth, and now the animal should move again in a line at right angles to the line connecting the two lights.

The following experiments carried out by Loeb and Northrop³⁰⁰ on the larvæ of the barnacle are perhaps the best proof for the validity of the Bunsen-Roscoe law for animal heliotropism.

These animals are small and can be obtained in large numbers. They were made to collect in the corner of a dish with a little sea water and were then sucked up into a pipette *cf.* Fig. 31, which was blackened with the exception of the opening. When such a pipette is put into a glass dish with parallel walls whose bottom is black (by putting paraffin blackened with lampblack at the bottom of the dish) the larvæ will flow out in a fine stream and swim when they are positively heliotropic in a straight line toward the source of light. They thus form a rather narrow white trail on the dark bottom and it is possible to measure the angle of this

trail with the line connecting the two lights. In this way in each observation the average trail of thousands of individuals is measured. By using one constant and one intermittent source of light and comparing the results with those obtained by two constant lights we can test the validity of the Bunsen-Roscoe law.

The method of the experiments was as follows: *abcd* (Fig. 31) is a square dish of optical glass with blackened bottom and containing a

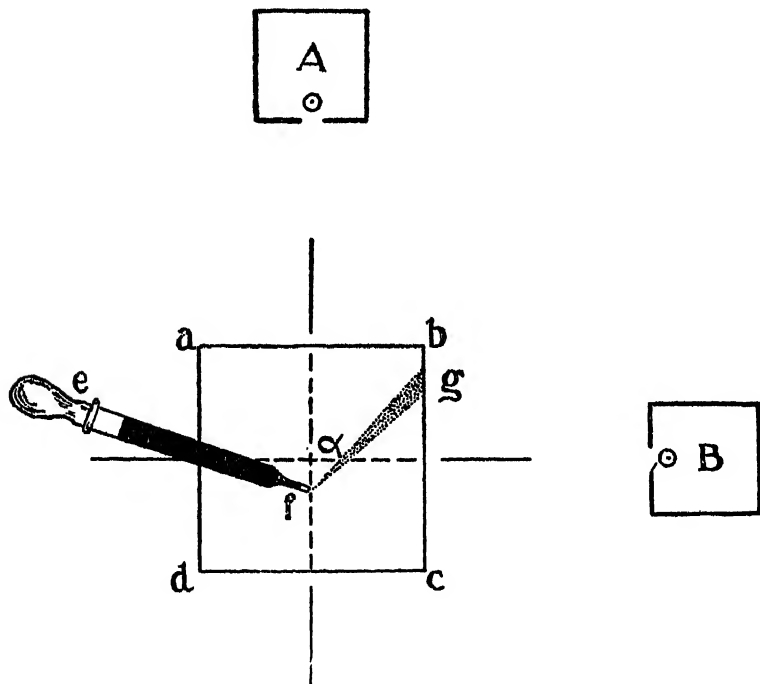


FIG. 31.—Method for the proof of the validity of Bunsen-Roscoe law for the positively heliotropic larvae of the barnacle. (After Loeb and Northrop.)

layer of sea water. *A* and *B* are two lights, the intensity of which is determined by a Lummer-Brodhun contrast photometer. In front of each light is a screen with a round hole permitting a beam of light to go to the dish. The lights and the dish *abcd* are so adjusted that the two beams of light striking the sides *ab* and *bc* at right angles cross each other in the middle of the dish. The light *A* is fixed while the light *B* is movable on an optical bench. The experiment is made in

a dark room and the lights *A* and *B* are enclosed in a box. At the beginning of the experiments the pipette is filled with a dense suspension of larvæ in sea water and then put with its point touching the bottom of the dish. The animals flow out in a fine stream which is narrow at the opening of the pipette and widens slightly, owing probably to the negative stereotropism of the animals.

A glass plate (Fig. 32) *hikl*, which has a strong red line *no* and a fine parallel line *pq* (cut with a diamond), is then put on the dish and so adjusted that *pq* is in the middle of the stream *fg* of the animals. Then the angle α which *pq* makes with the perpendicular from *A* on *ab* is measured. This perpendicular is marked in the form of a red line on the black base on which the glass vessel *abcd* stands. The angle α is measured with a goniometer. When the lights are equal in intensity α should be 45° ; if the two lights have different intensities and if *A* be the stronger light α should become smaller with increasing difference in intensity. The individual measurements vary comparatively little, as long as the difference in the intensity of the two lights is not too great; for this reason our observations do not go beyond a wider ratio of the two lights than 10:1, though 4:1 is perhaps the limit for good results. Table VII gives the results. *A* is always the stronger light. Each table is the average of from 40 to 60 individual observations, each being the average of the path of many thousands of animals.

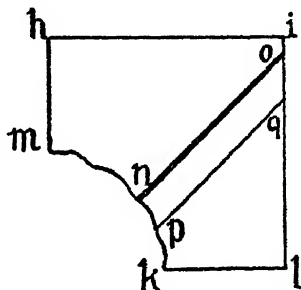


FIG. 32.

TABLE VII

| | Value of α for different ratios of intensities of the two lights | | | |
|--|---|------------|--------------|--------------|
| Ratio of the two lights..... | 1:1 | 2:1 | 4:1 | 10:1 |
| Value of α (direction of path)..... | 45.6° | 40° | 34.4° | 28.8° |

In the next series of experiments an opaque rotating disk with one sector cut out was placed before light *B*. In one set of experiments the sector cut out was 90° . The rate of rotation (by an electric motor) was 1,500 to 2,500 revolutions per minute. The other light was constant and its distance was chosen on the assumption of the validity of the Bunsen-Roscoe law for these cases. Thus when the two lights without sector were equal at a given distance of *A*, by putting 90° sector before

B, it was assumed that the ratio of effects would be the same as if, with constant light, *B* had been placed at the double distance and the ratio of intensities of the two lights had been 4:1. Going on such a calculation we should expect the same values for α as in Table VII.

As one sees from Table VIII, the observed values are slightly smaller but practically identical with the values obtained when the two lights are constant. The deviation is probably due to the well established fact that the photochemical efficiency of an intermittent light is a trifle less than that calculated on the basis of the Bunsen-Roscoe law.

TABLE VIII

| | Value of α when one light is intermittent (90° sector) and the other constant, and the efficiency of the two lights is calculated on the basis of the validity of the Bunsen-Roscoe photochemical law | | |
|------------------------------|--|-------|-------|
| Ratio of the two lights..... | 1:1 | 2:1 | 4:1 |
| Value of α | 44.2° | 38.3° | 34.1° |

We carried out some experiments with a sector of 141°. When the efficiency of both lights was equal on the assumption of the validity of the Bunsen-Roscoe law α was found to be 44.9° (instead of 45°), and for the ratio 2:1 α was found to be 38.8°. The values are, within the limits of error, identical with the values in Tables VII and VIII.³⁰⁶

Bradley M. Patten also showed that for the heliotropic reactions of the negatively heliotropic larva of the fly the law of Bunsen and Roscoe holds.

Photochemical processes have a very small temperature coefficient and it agrees with this that lowering of temperature within the limits compatible with the motility of animals does not affect the heliotropic response; on the contrary, we shall see that in certain crustaceans (*e.g.*, *Daphnia*) lowering of the temperature may enhance positive heliotropism.²⁹⁶

We must, therefore, conclude that the light produces in an eye or an element of the photosensitive skin a chemi-

cal reaction which results in the formation of a certain mass of a reaction product. This mass acts on the peripheral nerve endings and brings about an as yet unknown change in the brain elements with which these nerve endings are connected. This change in turn affects the tone or tension of the muscles with which the brain elements are connected. When the rate of photochemical reaction is the same in both eyes or in the photosensitive elements on both sides of the body, the change of tone in the symmetrical muscles of both sides of the body is the same and no change in the position or direction of motion of the organism should occur. If the rate of illumination is different in both eyes, differences in the relative tension of the symmetrical muscles occur, which make the motion to the source of light easy and in the opposite direction more difficult when the animal is positively heliotropic. For the negatively heliotropic animal the opposite effect will be brought about.

These experiments, therefore, show that the tropism theory not only allows us to predict the nature of the animal reactions but allows us to predict them quantitatively. Thus far the tropism theory is the only one which satisfies this demand of exact science.

The degree of directness with which a heliotropic animal goes to or from a source of light depends, aside from the degree of perfection of its locomotor apparatus, upon the intensity of the light and the relative sensitiveness of the animal. Animals which in strong light will move in approximately straight lines to or from the source of light may in weak light reach their goal in a more or less irregular zigzag line. This is easily understood. When

an animal by chance gets its median plane too far out of the direction of the rays of light (we assume them to be parallel), the rate of photochemical reaction will become different in both eyes. As soon as the difference between the photochemical reaction products in both eyes exceeds a certain limit the animal will automatically put its plane of symmetry again into the direction of the rays of light. The weaker the light and the less sensitive the animal, the longer it will take until this happens, and the greater the freedom of the animal to deviate from the straight line.

CHAPTER X

THE EFFECT OF RAPID CHANGES IN INTENSITY OF LIGHT

It may prove necessary to make a similar assumption for the effect of a constant illumination as was made by Nernst for the theory of the action of galvanic currents, namely that there are two antagonistic processes going on, one being the photochemical effect of light and the second either a process of diffusion of the substances formed or a chemical reaction of the opposite character as that caused by the action of the light. Many animals which are oriented by constant illumination react by a quick, jerky movement when the intensity of light is either rapidly increased or diminished. In this case the effect is determined by the rapidity of the change in the intensity, $\frac{di}{dt}$, and not by the product of intensity into duration of illumination, *it*.²⁹⁷ These twitching or jerking effects caused by a rapidly changing intensity of light are comparable to the twitching brought about in a muscle by a rapid increase or decrease in the intensity of a current. The writer described such reactions first for tube worms like *Serpula*, which withdraws suddenly into its tube when a shadow passes over it or when the intensity of light is suddenly diminished in some other way. The anthropomorphists, of course, declare this reaction to be induced by the instinctive fear of an enemy, oblivious of the fact that if they were consistent they would have to give the same explanation for the twitching of a muscle upon rapid changes in the intensity of a current. The

problem to be solved is in both cases a purely physico-chemical one. It was also found that the motions of certain animals stop when they come suddenly from strong light into weak light. This was observed in planarians which as a consequence collect in greater density in spots

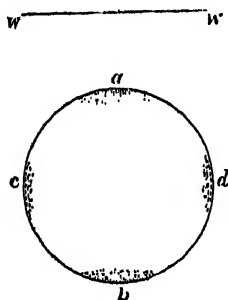


FIG. 33.—Difference in place of gathering between heliotropic animals and animals which come to rest when reaching a relative minimum in the intensity of light. In a circular vessel $a\ b\ c\ d$ and $W\ W$ representing the window, positively heliotropic animals will collect at a , negatively heliotropic animals at b , and animals which come to rest where the intensity of light is a relative minimum at c and d .

of the space where the intensity of light is a relative minimum.²⁹¹ The difference in the conduct of heliotropic organisms like *Daphnia* which go to or from the light and animals like planarians which come to rest where the intensity of light is a relative minimum can be demonstrated by putting them into a circular vessel (Fig. 33). The positively heliotropic animals collect at a , the negatively heliotropic at b , while the planarians collect at c and d where the intensity of light is a minimum. Reactions determined by the value $\frac{di}{dt}$ do not lead to phenomena of orientation, though such (improperly called) "fright reactions"^a occur in many helio-

tropic animals; they may lead, however, to collections of animals.

Jennings has maintained that all reactions of unicellular organisms are due to "fright" or "avoiding reac-

^a The reader should notice the difference in the treatment of animal conduct from the point of view of the physicist and of the introspective psychologist. What the physicist expresses correctly by the term $\frac{di}{dt}$ the anthropomorphic biologist explains in terms of human analogy as "avoiding reaction" or "fright reaction," a term which not only assumes the existence of sensations without any adequate proof, but removes the problem from the field of quantitative experimentation.

tions," and it seems as if at one time he even intended to deny the existence of tropisms and to maintain that all animals were influenced only by rapidly changing intensities of light. It is needless to discuss such an idea (which he probably no longer holds) in view of the contents of the preceding chapters. He seems, however, to cling to it as far as asymmetrical unicellular organisms are concerned. When moving about *Paramæcia* often reverse the direction of their progressive motion for a moment, but then do not return in the old direction, moving sidewise, on account of the asymmetry in the arrangement of their cilia. Jennings is probably right in assuming that this factor can lead to collections of such infusorians, since it may prevent their leaving a drop and going into the surrounding medium. When, *e.g.*, at the boundary of the two media such a reversal of the action of the cilia occurs, the organisms are prevented from crossing from one medium into the other.

But Jennings goes too far in this attempt, when he tries to explain the heliotropic reactions of certain unicellular organisms, *e.g.*, *Euglena*, in this way. He maintains ²⁵³ that unicellular organisms like *Euglena* go to the light on account of shock movements produced by the shading of the photosensitive region of the animal. *Euglena* moves with a constant rotation around its longitudinal axis and Jennings assumes that in a certain phase of the rotation a photosensitive element (the eye spot) of the organism is shaded. This he thinks causes a shock movement, whereby the animal is swerved to the light again during the next half of the spiral revolution, and so on. Similarly in negatively heliotropic *Euglena* the swerving away from the light is, according to Jennings, the shock movement caused by the increased illumination

of the photosensitive end of the animal produced by swerving toward the light during the previous half of the spiral revolution. Bancroft²¹ showed that Jennings's theory was based upon incomplete facts.

According to Jennings's view positive heliotropism is conditioned by and should be accompanied by shock movements produced by sudden shading (= shading reaction), and negative heliotropism should always be accompanied by shock movements produced by sudden illumination.

It has been found, however, that this usual association of shock movements with tropism is not a necessary one, but that it can be destroyed if the proper means be taken. Consequently the view that the heliotropic swerving is a shock movement must fall.

When *Euglenæ* from Culture B were placed in the rays of the arc light, at a distance of four or five feet from the light, they were strongly positively heliotropic and gave the shading reaction. When, however, they were gradually brought nearer to the light a point was reached at which the heliotropism disappeared but the shading reaction persisted. When moved still closer to the light they became negatively heliotropic but still without any change of the shading reaction. When moved still closer to the light, there was a short time when no shock movements could be obtained, but soon the illumination reaction appeared. At the same time the negative heliotropism became more prompt and precise. Finally, when the light was still further increased and allowed to act for a considerable time, even the illumination reactions frequently disappeared completely, and a most pronounced and compelling negative heliotropism held full sway. . . .

It is very evident, then, that the invariable correlation of positive heliotropism with the shading reaction, which is required by Jennings's theory, does not exist. Both kinds of heliotropism may be associated with either the shading or the illumination reaction. Accordingly, it must be concluded that the heliotropic mechanism does not depend upon the mechanism for the shock movements, but that the two mechanisms are independent.²¹

The simplest method of determining whether or not the orientation of flagellates depends upon rapid changes in intensity of light or upon constant illumination can be furnished with the aid of intermittent light. We know that a striped muscle contracts only when a current is

made or broken, but not while the constant current lasts. Hence a rapidly alternating current throws the muscle into tetanus, while the constant current has no effect. If it is the rapid change in the intensity of light which causes the swimming of a positively heliotropic *Euglena* to the light, an intermittent light, of a sufficient number of alternations per second, should be much more efficient than a constant light; while in case the positive heliotropism is determined by constant illumination, this should not be the case and the Bunsen-Roscoe law should hold.

Mast³⁴⁸ has recently published experiments on the relative efficiency of the various parts of the spectrum by a method based on the assumption of the validity of the Bunsen-Roscoe law for the heliotropic orientation of these organisms. If his assumption^b is correct, it contradicts the theory which Jennings and Mast have defended now for more than fifteen years; if his assumption is wrong, his experiments on the relative efficiency of various parts of the spectrum cannot be correct. Since, however, Mast's results with this method coincide with those by Loeb and Wasteneys³¹² obtained by a direct method, it is very probable that the law of Bunsen and Roscoe holds for the heliotropic reactions of *Euglena* and unicellular flagellates in general, and, if this is true, the heliotropic reactions of unicellular algæ (*Euglena* included) are determined by light of constant intensity.

^b He does not seem to have noticed that his method was based on this assumption.

CHAPTER XI

THE RELATIVE HELIOTROPIC EFFICIENCY OF LIGHT OF DIFFERENT WAVE LENGTHS

1. The validity of the Bunsen-Rosecoe law for the heliotropic reactions of animals and plants leaves no doubt that these reactions are determined by the rate of photochemical processes. Heliotropic reactions depend, however, not only upon the intensity but also upon the wave length of light. Photochemistry shows that the most efficient wave length varies with the nature of the photochemical substance and that comparatively slight changes in the constitution of a molecule may bring about considerable changes in the relative efficiency of different wave lengths. The search for differences in the heliotropic effect of different wave lengths can be of service in detecting the nature of the photochemical substances responsible for heliotropic reactions.

The investigations on the relative heliotropic efficiency of different wave lengths have generally been undertaken for a different purpose, namely, to get information concerning the color sensations of animals. Graber gave it as the result of his observations that all animals which were fond of light were also fond of blue, and animals which were fond of dark were also fond of red.¹⁸⁰ He put animals into a box half of which was covered with transparent glass and half with an opaque object, and then counted the relative numbers of organisms in both halves of the box. He then replaced these screens by colored glasses and obtained the above-mentioned result. The

writer showed that the animals are neither fond of blue nor of red but are oriented by the light in the same way as are plants, and the statement that animals which were "fond" of light also were "fond" of blue and those which were "fond" of "dark" were "fond" of red the writer explained in a simpler way, namely that the light filtered through red glass had a smaller orienting effect than the light filtered through blue glass.²⁸⁷ Hence red glass acted like an opaque, blue glass like a transparent screen. This had already been known to be true for the heliotropic reactions of plants for which Sachs had shown that they occur behind a blue glass in the same way as behind common window glass, while behind red glass heliotropic reactions do not occur at all or occur very slowly as if the light were weak. The writer was able to show that the same is true for animals.²⁸⁷ When positively heliotropic animals are put into a box covered with blue glass they go as rapidly to the window side as when the box is uncovered; while when it is covered with red glass the animals will go to the window but more slowly and irregularly. Behind a red screen they behave therefore as if they were exposed to weak light.

Blue glass is permeable not only for blue but also for rays which produce the sensation of green. Paul Bert²⁸ had already made experiments with positively heliotropic *Daphnia* in a solar spectrum and found that the animals "accouraient beaucoup plus rapidement au jaune ou au vert qu' à toute autre couleur."^a Bert concluded from this that to the eye of a *Daphnia* those parts of the spectrum appear brightest which also appear brightest to the human eye. Bert's aim was to find out whether the sensa-

^a This method of ascertaining the most efficient part of the spectrum is not reliable and has been replaced by other methods.

tions caused by light in lower animals are the same as those caused in a human being. But even if the relative efficiency of the various parts of the spectrum were the same for the sensations of brightness in human beings and for the heliotropic reactions in lower animals, it would not prove that the latter also have sensations of brightness. For we have no guarantee that the heliotropic reactions of lower animals are due to or accompanied by sensations of brightness. If the yellow-green rays are the most efficient in causing heliotropic reactions in an organism, it suggests only that in such an organism the photosensitive substance responsible for the heliotropic response is most easily decomposed by the yellow-green part of the spectrum.

A similar error of reasoning as that by Bert has recently been made by Hess. Hess corroborated what the writer had already pointed out, that the red rays of the visible solar spectrum are the least efficient, and he found, moreover, as Bert had found for *Daphnia*, that for the heliotropic reactions of a number of animals, from the fishes down, the yellow-green region of the solar spectrum is the most efficient. Now it happens that to a totally color blind human being, to whom the different parts of the solar spectrum appear only as shades of gray, the region $\lambda = 540 \mu\mu$ in the yellow-green appears to be the brightest; while the red part of the spectrum gives a very faint sensation of brightness. From this similarity or apparent identity between the relative effects of different wave lengths upon the heliotropic effects of certain lower animals and upon the sensations of brightness of a totally color blind human being, Hess draws the conclusion that these animals are totally color blind. In our opinion the only conclusion which Hess has a right to

draw is that the photosensitive substance which causes sensations of brightness in the eye of the color blind human being is either identical with or is affected in a similar way by light waves as is the substance giving rise to heliotropic reactions in certain animals. This assumption is entirely adequate and harmonizes better with the facts than the assumption made by Hess. The substance responsible for the sensations of brightness in the eyes of the totally color blind human being is visual purple which is bleached most rapidly by light of $\lambda = 540 \mu\mu$. That our objection is justified is proved by the experiments of v. Frisch on bees.

v. Frisch¹⁶⁸ has shown by very ingenious and careful experiments that bees can be trained to discriminate between blue and yellow but not between different shades of gray. On a table were put square cardboards of different shades of gray and among them one blue piece of cardboard. On each gray square was put a watch crystal containing water, while the watch crystal on the blue contained sugar water. The bees of an observation hive visiting the sugar crystal were marked with a fine paint brush. After a sufficient period of training it was found that the marked bees always went directly to those crystals which were on a blue piece of cardboard, whether they contained sugar water or pure water; and when there was no sugar water on the blue cardboard they alighted on any blue object, *e.g.*, a blue pencil. The crystals and cardboard pieces were always renewed in different tests, to avoid any influence of odor. It was never possible to train the bees to select a piece of cardboard with a definite shade of gray among cardboards of different shades of gray.

Hess had shown that for the heliotropic reaction of

bees the yellowish-green part of the spectrum is most efficient, and he concluded that bees are totally color blind. To a totally color blind person the blue cardboard appears only like a shade of gray, and such a person is unable to learn to discriminate between a blue and gray piece of cardboard. v. Frisch's experiments support the conclusion that it is unjustifiable to use experiments on heliotropism to draw conclusions concerning light or color sensations. v. Frisch and Kupelwieser¹⁸⁹ have also demonstrated selective effects of different light waves for *Daphnia* which differ from those found for the eye of a totally color blind person, and their observations have been confirmed by Ewald.¹⁴⁷

2. Hess's conclusions are in conflict with another group of facts. For many plants the blue region of the solar spectrum is the most efficient. Hess is, therefore, compelled to conclude that the heliotropism of such plants is different from that of animals, since it would seem preposterous to assume that swarmspores of plants should go to the light because they have sensations of color. He, therefore, assumes that plants are heliotropic—in the sense of the mechanistic theory—but that positively heliotropic animals go to the light on account of their love for brightness which is exactly the old viewpoint of Graber. It can be shown, however, that the difference in the heliotropism of animals and plants, which Hess assumes, is contrary to the facts, since there are heliotropic animals for which the blue rays are the most efficient, as for most plants; and there are green algæ for which the yellowish-green rays are most efficient, as for animals.

For many if not most plants the blue rays are the most efficient for inducing heliotropic curvatures. Blaauw⁴⁷ proved this in the following way: He exposed a row of

seedlings of *Avena* to a carbon arc spectrum for a certain time. The seedlings were then placed in the dark and after the proper time it was ascertained which part of the spectrum had induced heliotropic curvatures. By varying the duration of time of exposure to the spectrum it was found that with a minimal time of exposure only certain blue rays, namely, those of a wave length of $478\ \mu\mu$ caused heliotropic bending, while with longer exposure longer waves also became efficient. In this way the minimum duration of exposure for various parts of the spectrum was ascertained. Table IX gives his results.

TABLE IX

| Duration of illumination, in seconds | Location of threshold in the spectrum, in $\mu\mu$ |
|--------------------------------------|---|
| 6.300 | 534 $\mu\mu$ |
| 1,200 | 510 $\mu\mu$ |
| 120 | 499 $\mu\mu$ |
| 15 | 491 $\mu\mu$ |
| 5 | 487 $\mu\mu$ |
| 4 | 478 $\mu\mu$ |
| 3 | 466 $\mu\mu$ |
| 4 | 466 $\mu\mu$ |
| 6 | 448 $\mu\mu$ |

The red and yellow parts of the spectrum were ineffective for the intensity and time limits used and the optimum of efficiency was in the blue, in the region between 466 and $478\ \mu\mu$.

A shorter series of experiments was made on the fruit bearers of *Phycomyces*, with the following results:

44 to 47 per cent. of the *Phycomyces* showed heliotropic curvatures

after 192 seconds of illumination at $615\ \mu\mu$
 after 192 seconds of illumination at $550\ \mu\mu$
 after 16 seconds of illumination at $495\ \mu\mu$
 after 32 seconds of illumination at $450\ \mu\mu$
 after 64 seconds of illumination at $420\ \mu\mu$

The number of experiments was limited but they indicate an optimum between 495 and 450 $\mu\mu$, in this respect agreeing with the results on *Avena*.

The fact then exists that for the heliotropic reactions of certain plants the blue rays are most efficient, while for the heliotropic reactions of a number of animals the yellowish-green rays are most efficient. But this statement cannot be generalized.

Loeb and Wasteneys determined the most efficient wave length of light for various lower organisms with the result that there are heliotropic animals for which the blue rays are as efficient as they are for plants; and that for different unicellular green organisms the optimum lies in different parts of the spectrum. They found, by a method similar to that used by Blaauw, that for the heliotropic curvature of the animal *Eudendrium* the most efficient part of the spectrum lies in the blue $\lambda =$ approximately 473 $\mu\mu$.³¹¹ The same was found by them for the larvæ of the marine worm *Arenicola*.

On the other hand, on investigation of two closely related forms of green flagellates, *Euglena* and *Chlamydomonas*, it was found³¹¹ that they behave differently. For *Euglena viridis* the blue rays $\lambda = 470$ to 480 $\mu\mu$ are especially efficient, while for *Chlamydomonas pisiformis* the most efficient part was in the region of $\lambda = 534 \mu\mu$, in the yellowish-green.^b For another green algæ, *Pandorina*, Loeb and Maxwell had already found the greatest efficiency in the greenish-yellow.

^b This would lead us, on the basis of the reasoning of Hess, to the conclusion that the unicellular plant *Chlamydomonas* has sensations of brightness, suffers from total color blindness (although it has no eyes), that it is not heliotropic, and that it is an animal; while its unicellular cousin, *Euglena*, has a highly developed color sense, has no sensations of brightness, is heliotropic, and is a plant.

The method used for these experiments by Loeb and Wasteneys is as follows:

A carbon arc spectrum, about from 18 to 23 cm. wide, was thrown on a black screen SS (see Fig. 34) with two slits a and b in the two different parts of the spectrum which were to be compared in regard to their heliotropic

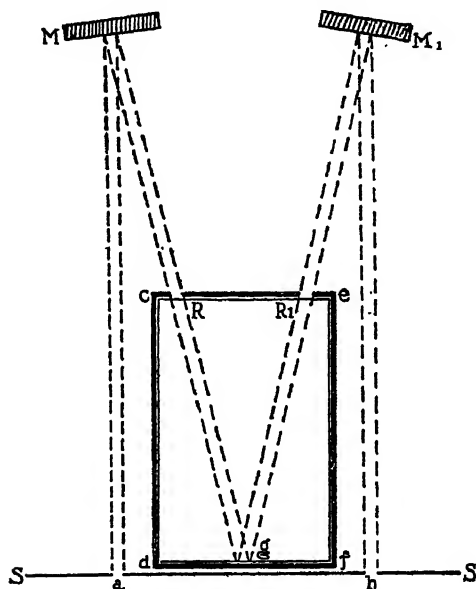


FIG. 34.—Method of determining the relative heliotropic efficiency of two different parts of the spectrum. (After Loeb and Wasteneys.)

efficiency. The two beams of light passing through the slits are reflected by the two mirrors M and M_1 into the square glass trough $cdfe$ in such a way as to strike the same region g of the back wall of the trough. The glass trough is surrounded by black paper except at R and R_1 , where the two beams of light enter from the mirrors. Before the experiment begins, all the organisms are collected in the spot g with the aid of an incandescent lamp. As

soon as the spectrum is turned on, these organisms are simultaneously exposed to two different beams of light which come from the two mirrors M and M_1 . When one type of light, *e.g.*, that from M , is much more efficient than the other coming from M_1 , practically all the organisms are oriented by the light from M and move toward this mirror, collecting in the region R . When the relative efficiency of the two types of light is almost equal, the organisms move in almost equal numbers to R and R_1 . By using as a standard of comparison the same region of the spectrum and successively altering the position of the other slit in the spectrum, we were able to ascertain with accuracy the relative efficiency of the different parts of the spectrum for the two forms of organisms. When the two parts of the spectrum which are to be compared are very close to each other, it is necessary to deflect the beams with the aid of deflecting prisms, before they reach the two mirrors.³¹¹

Experiments on the newly hatched larvæ of *Arenicola*, a marine worm, showed that the most efficient part of the spectrum was in the bluish-green of about $\lambda = 495 \mu\mu$, while for the larvæ of *Balanus eburneus* the most efficient part of the spectrum was found by Loeb and Maxwell, by Hess, and by Loeb and Wasteneys in the region of yellow and yellowish-green.³¹¹

Mast³⁴⁸ made similar experiments on these organisms with a method in which the organisms were exposed to two beams of light of different wave length crossing each other at right angles. One light was kept constant while the other was made intermittent by a disk with a sector cut out rotating in front of the light. The size of the sector was varied until the organisms moved at an angle of 45° to the two beams. When this happened the heliotropic

WAVE LENGTH

TABLE X I
RELATIVE STIMULATING EFFICIENCY OF DIFFERENT REGIONS IN THE SPECTRUM; REDUCED TO THE SAME MAGNITUDE AND AVERAGED SO THAT THEY CAN BE READILY COMPARED. MAXIMUM IN HEAVY TYPE

| Wave length in μ | <i>Euglena viridis</i> | | <i>Phacus triqueter</i> | <i>Trache- monas euchlora</i> | <i>Gonium</i> | <i>Areni- cola latens</i> | <i>Lum- bricus</i> | <i>Chlamy- domonas</i> | <i>Blowfly</i> | <i>Pando- rina</i> | | <i>Spondy- lomorpha</i> | |
|----------------------|------------------------|----------|-------------------------|---------------------------------------|---------------|-----------------------------------|------------------------|----------------------------|----------------|------------------------|----------|-----------------------------|-------|
| | Negative | Positive | | | | | | | | Negative | Positive | | |
| Blue. | 422.4 | 2.50 | 4.29 | 7.22 | 3.55 | 11.90 | 5.49 | 11.84 | 6.26 | 14.83 | 0.24 | 1.32 | 5.62 |
| | 432.6 | 4.09 | 7.69 | 12.21 | 15.81 | 16.60 | 14.01 | 9.72 | 16.17 | 18.91 | 2.65 | 3.57 | 7.62 |
| | 442.8 | 10.41 | 12.70 | 16.49 | 16.60 | 20.10 | 16.63 | 20.14 | 18.91 | 18.91 | (?) | 4.89 | 10.85 |
| | 452.9 | 13.24 | 16.66 | 19.02 | 18.59 | 20.03 | 20.24 | 20.14 | 18.91 | 18.91 | 16.17 | 7.86 | 13.87 |
| | 463.1 | 14.86 | 18.30 | 20.07 | 20.03 | 17.50 | 20.24 | 20.14 | 18.91 | 18.91 | 16.17 | 9.15 | 16.35 |
| Green. | 483.4 | 20.40 | 20.86 | 20.07 | 20.03 | 17.50 | 20.24 | 20.14 | 18.91 | 18.91 | 16.17 | 9.15 | 16.35 |
| | 493.6 | 14.46 | 18.30 | 16.47 | 17.64 | 19.45 | 17.34 | 18.23 | 17.96 | 19.39 | 13.39 | 12.51 | 16.60 |
| | 503.7 | 8.23 | 9.46 | 8.42 | 10.21 | 13.37 | 11.37 | 7.84 | 19.88 | 20.88 | 15.15 | 15.15 | 19.30 |
| | 513.8 | 3.06 | 3.13 | 1.98 | 5.16 | 5.76 | 6.58 | 2.34 | 15.70 | 19.50 | 16.21 | 18.78 | 19.62 |
| | 524.0 | 1.08 | 1.00 | 0.30? | 1.05 | 2.71 | 2.67 | 8.01 | 13.33 | 16.21 | 16.21 | 20.10 | 19.27 |
| Yellow. | 534.1 | | | | | 0.29 | 7.99 | | | | 13.33 | 18.51 | 20.17 |
| | 544.3 | | | | | | | | | | 8.41 | 17.67 | 18.20 |
| | 554.4 | | | | | | | | | | 5.71 | 14.25 | 12.95 |
| | 564.5 | | | | | | | | | | 3.42 | 0.42 | 7.00 |
| | 574.6 | | | | | | | | | | 1.72 | 6.57 | 5.00 |
| Red. | 584.8 | | | | | | | | | | 2.95 | 3.39 | 2.95 |
| | 594.9 | | | | | | | | | | 0.51 | 1.53 | 2.30 |
| | 605.0 | | | | | | | | | | 0.40 | 0.60 | 1.70 |
| | 615.2 | | | | | | | | | | 0.22 | 1.35 | 1.35 |
| | 625.3 | | | | | | | | | | 0.19 | 0.09 | 1.00 |
| Red. | 635.4 | | | | | | | | | | | | 1.00 |
| | 645.6 | | | | | | | | | | | | 0.42 |

efficiency of the two beams was considered equal. Mast's results, which are given in Table X, agree with those of Loeb and Wasteneys.

The error which Hess makes is of epistemological interest inasmuch as it shows the danger of false analogy. The real analogy for heliotropic reactions are forced movements and other tropisms, *e.g.*, galvanotropism or geotropism. Since forced movements (*e.g.*, in Mènière's disease) and galvanotropic reactions caused by a constant current through our head are not determined or accompanied by special sensations, the same may be true in regard to heliotropic reactions. This is not an idle assumption, since we know that the contraction of the iris of our eye under the influence of light is not accompanied by any sensation of brightness or color and such contractions occur also under the influence of light when the iris is excised. Hess ignores not only this analogy, but the whole existence of forced movements and of other tropisms, and he uses the color and light sensations of human beings, who are not heliotropic, to explain heliotropism in animals about whose sensations we know nothing. He fails to see that by this false analogy he dodges the real problem of heliotropism, namely, why the tension of symmetrical muscles changes upon one-sided illumination of an animal. For the explanation of this problem, we find assistance in the field of forced movements and of galvanotropism and of geotropism, but not in the behavior of totally color blind human individuals who show no trace of heliotropism.

The adoption of the false analogy between visual sensations and heliotropism makes it impossible for Hess to admit that bees should be heliotropic and at the same time be able to discriminate between blue and gray; while

if we take cognizance of the analogy between heliotropism and the other tropisms we realize that the heliotropism of the bees and their reactions to blue are separate and independent phenomena, which need not be mutually exclusive and which in all probability depend upon different parts of the brain. When in certain cases the relative heliotropic efficiency of the various parts of the spectrum is identical with the curve for its apparent relative brightness to a totally color blind person, we may conclude that the photosensitive substances responsible for the two groups of phenomena behave similarly or may even be identical, but not that the sensations of brightness of the color blind and the heliotropic reactions of insects are identical or analogous phenomena.

Many mutants of *Drosophila* differ in regard to the pigments of the eye. It was natural to raise the question whether or not such hereditary variations of pigmentation of the eye influence the reaction of the flies to monochromatic light. McEwen investigated this possibility with the following result: "Colored lights which may be conveniently described as violet, green and red, are effective in the order named upon the insects whose eye color is lighter than the red eye of the wild fly. In the case of wild flies, and flies whose eyes are of a still darker shade called sepia, red is more effective than green" (McEwen⁵⁴⁹).

CHAPTER XII

CHANGE IN THE SENSE OF HELIOTROPISM

WE have stated that while in a positively heliotropic animal a one-sided illumination increases the tension of the muscles which turn the animal toward the source of light, in the negatively heliotropic animal the one-sided illumination must result in the opposite effect, namely, in a diminution of tension in the same muscles. As a consequence, the negatively heliotropic animal can turn more easily away from the light than toward the light.

Groom and Loeb¹⁸³ noticed that the larvæ of the barnacle upon hatching go directly to the light and gather at the light side of a dish, but that sooner or later their positive heliotropism may give way to an equally pronounced negative heliotropism. The stronger the light the more rapidly the larvæ are transformed into negatively heliotropic organisms. Later the reversibility of the sense of heliotropism was observed and studied in a number of organisms.²⁹¹ In a summary of the subject³⁰⁰ (p. 470) the writer pointed out that this reversion was due either to a modification of photochemical processes or to an effect upon the nervous system. That an influence on the nervous system can indeed bring about a change in the sign of the reaction is very strikingly demonstrated in the following observation of A. R. Moore on starfish.⁵²⁵

Ordinarily, when a starfish which is moving in an aquarium is touched, it stops immediately and clings tenaciously to the surface of the vessel with its tube feet, so that it is impossible to remove the animal without injury to the tube feet. This normal response to sudden contact can be completely reversed by the administration of strychnine, so that when touched the animal loosens its hold on the bottom completely.

The starfish poisoned with strychnine upon sudden touch withdraws all the tube feet, so that it can be moved about like an inert object. For this purpose 1 or 2 c.c. of a 0.5 per cent. solution of strychnine sulfate were injected into a starfish of medium size.

If the stretching out of the tube feet is due to an increase in the tone of the ring muscles (and a decrease in the tension of the longitudinal muscles) the drawing in is due to an increase in the tone of the longitudinal muscles of the tube feet. We therefore see that the same "stimulus," namely, a sudden touch, which causes one set of muscles to contract in a normal animal causes the antagonists of these muscles to contract in an animal poisoned with strychnine. We shall see that a number of cases of reversal of heliotropism may well find their explanation on this basis. On the other hand, the phenomena of solarization known in photography indicate that the sign of heliotropic response may also be changed by an excessive action of light on the photochemical substance. This effect, of course, may in the last analysis also result in an influence upon the central nervous system, such as that brought about by strychnine in Moore's experiment. We will now consider some cases more in detail.

The writer found²⁹⁶ that certain fresh water crustaceans, namely Californian species of *Daphnia*, copepods, and *Gammarus* when indifferent to light can be made intensely positively heliotropic by adding some acid to the fresh water, especially the weak acid CO_2 . When carbonated water (or beer) to the extent of about 5 or 10 c.c. is slowly and carefully added to 50 c.c. of fresh water containing these *Daphnia*, the animals will become intensely positive and will collect in a dense cluster on the

window side of the dish. Stronger acids act in the same way but the animals are liable to die quickly. Esters, *e.g.*, ethylacetate, act also like acids and the addition of 1 c.c. of a grammolecular solution of ethylacetate to 50 c.c. fresh water also makes all the organisms positively heliotropic. Alcohols act in the same way. In the case of *Gammarus* the positive heliotropism lasts only a few seconds, while in *Daphnia* it lasts from 10 to 50 minutes and can be renewed by the further careful addition of some CO₂. The following table gives the minimal concentration of various acids and alcohols for the production of positive heliotropism in certain California species of fresh water copepods, and *Daphnia*:

| | For Copepods | For <i>Daphnia</i> |
|----------------------------|--------------|--------------------|
| Formic acid | 0.006 N | |
| Acetic acid | 0.006 N | |
| Propionic acid | 0.005 N | |
| Butyric acid | 0.004 N | |
| Valerianic acid | 0.004 N | |
| Capronic acid | 0.002 N | 0.6 N |
| Ethyl alcohol | 0.19 N | 0.2 N |
| Propyl alcohol | 0.054 N | 0.05 to 0.1 N |
| Normal butyl alcohol | 0.019 N | |
| Isobutyl alcohol | | 0.04 N |
| Amyl alcohol | 0.011 N | |

As far as alcohols are concerned each higher alcohol is about three times as efficient as the previous one, with the exception of amyl alcohol. This order of relative efficiency is also characteristic for the surface tension effects of these alcohols.²⁹⁹

It was of importance to find means of making these organisms negatively heliotropic. Moore³⁶⁸ found that caffeine makes the heliotropically indifferent fresh water crustacean *Diaptomus* intensely negatively heliotropic. It required the addition of 1.2 c.c. of a 1 per cent. solution of caffeine to 50 c.c. of water to bring about this intense

negativation. In two minutes all the animals are collected in a dense cluster on the negative side which lasts for about 35 minutes. A weak negative collection could also be obtained by adding 0.1 c.c. of a 0.5 per cent. solution of strychnine nitrate. Moore found that if the *Diaptomus* were first made positively phototropic by the addition of alcohol or acids, it was impossible to alter their response by the action of caffeine, strychnine, or atropine. On the other hand, animals which had formed a negative collection under the influence of caffeine if treated with carbonated water at once changed their response and swimming to the light side of the dish formed a positive gathering.

What causes these effects? The fact that alcohols make the organisms positively heliotropic suggested the possibility of a "narcotic" effect; the writer found, however, that narcosis requires a concentration of alcohols three times as high as the one required to produce positive heliotropism. He tried the effect of temperature on the reversal of the sign of heliotropism in *Daphnia* and found that lowering of the temperature enhanced the effect of acids in making the animals positive.²⁹⁶

The writer had found previously that in marine crustaceans and in larvæ of a marine annelid, *Polygordius*, the sense of heliotropism can be reversed by changes of temperature as well as by changes in the osmotic pressure of the sea water.²⁹¹ Increase in the osmotic pressure of sea water (by adding about 1 gm. of NaCl or its osmotic equivalent of other substances to 100 c.c. of sea water) made the negative animals positively heliotropic, and lowering of the concentration by adding 30 to 60 c.c. distilled water to 100 c.c. sea water made positive organisms negative. Negative larvæ of *Polygordius* or negative

marine copepods could be made positive by lowering the temperature, and positive larvæ could be made negative by slowly raising the temperature. Since in the latter case the animals suffered from the high temperature the results were not so striking as in the case of the positivating effect of lowering the temperature. The same effect of the concentration of sea water and of temperatures was observed by Ewald for the larvæ of *Balanus perforatus*. He found, moreover, the interesting fact that a change of the ratio $\frac{Na}{Mg}$ in the sea water affected the sign of heliotropism of barnacle larvæ. An increase of Na made them more positive, an increase in Mg more negative.¹⁴⁴

The larvæ of *Porthesia* are strongly positively heliotropic before they have eaten, while they lose their heliotropism almost completely after they have eaten.²⁸⁷ The writer observed that male and female winged ants are strongly positively heliotropic but as soon as they lose their wings their heliotropism ceases.²⁸⁷ McEwen⁵⁴⁹ has found that when *Drosophila* is deprived of its wings its heliotropism ceases.

Holmes found that terrestrial amphipods are positively, while the aquatic amphipods are negatively heliotropic. By putting a terrestrial amphipod into water it became negatively heliotropic.²²⁵

That a reversal in the sense of heliotropism may be due to a nervous effect is suggested by an observation by Miss Towle⁴⁸⁵ that a certain ostracod, *Cypridopsis*, can be made positively heliotropic by mechanical shock, and the writer noticed that indifferent fresh water *Gammarus* can be made negatively heliotropic by shaking them. In both cases the heliotropism lasts only a short time.

The attempt to explain all these reversals on the assumption of a change in the central nervous system meets with the difficulty that such reversals occur also in unicellular organisms which have no central nervous system. Thus the writer observed that *Volvox*, which occurred in the same ponds in California from where *Daphnia* came, could also be made positive by CO_2 .²⁹⁶ In swarmspores of algæ reversals of heliotropism are a common phenomenon. While these unicellular organisms have no central nervous system they may have synapses such as exist between different neuria of metazoa. The writer is not sufficiently familiar with the behavior of synapses in higher animals to suggest that this condition is responsible for the changes in the sense of heliotropism.

We may finally discuss briefly a possible solarization effect. The writer found that it is possible to make animals generally negatively heliotropic with the aid of ultraviolet light.²⁹⁶ If once rendered negative such animals will be negative not only to ultraviolet rays but also to the light of an incandescent lamp. A. R. Moore³⁰⁶ found that the ultraviolet rays having such an effect have a wave length shorter than 3341 Å.U. Oltmanns had observed that *Phycomyces* is positively heliotropic in weak light, indifferent in somewhat stronger light, and negatively heliotropic in still stronger light. Blaauw found that when the illumination was strong the seedlings of *Avena* became negatively heliotropic.⁴⁷ He suggests the analogy with solarization effects in photography. The discovery of photodynamic effects by v. Tappeiner⁴⁷⁷ adds to the possibilities which should be considered in this connection.

While *Drosophila* is usually positively heliotropic, McEwen has recently described a mutant of this species

which is not heliotropic. This lack of heliotropic response is linked with a peculiar color—"tan"—by which the mutant is characterized. The character "tan" is sex linked. The daughters inherit the factor for the character from their fathers but do not show the character, while the sons inherit the factor from their mothers and do show the character. The lack of heliotropic reaction in this mutant is apparently not due to any structural defect in the eye (McEwen⁵⁴⁹).

Keeping successive generations of flies in the dark does not influence their heliotropism. F. Payne^{550, 551} raised sixty-nine successive generations of *Drosophila* in the dark, but the reaction of the insects to light (as well as their eyes) remained entirely normal.

CHAPTER XIII

GEOTROPISM

1. When the stem of certain plants is placed in a horizontal position, the apex grows vertically upward and the root downward. The downward growth of the root is called positive, the upward growth of the apex negative geotropism. The writer has observed a similar phenomenon in a hydroid, *Antennularia antennina*^{294, 300} and his observations were confirmed by Miss Stevens.⁵⁵³ Animals as well as plants, therefore, show the phenomenon of geotropism.

These phenomena have given rise to a strange discussion, namely: What constitutes the "stimulus" in the case of geotropism? When a galvanic current is sent through a motor nerve the muscle answers with a contraction only when the current is made or broken, but not while a constant current is flowing through the nerve. The older physiologists were not able to form a mental picture of what happened in this case, and they cut the knot by invoking a verbalism, namely by calling the making or breaking of a current a "stimulus." This perhaps innocent verbalism then led to the less harmless dogma that only a rapid change could act as a "stimulus." Thus Jennings²⁵³ and Mast³⁴⁶ took it for granted that phenomena of orientation by light could only be produced by rapid changes in the intensity of light and not by constant illumination, since they had the *a priori* conviction that only a rapid change in the intensity of a galvanic current or of light is a "stimulus." The same diffi-

culty arose in regard to the action of gravity upon orientation, since it was contrary to the definition of a "stimulus" that the mere permanent lying in a horizontal position should cause the apex of a stem to bend upward.

All these difficulties disappear if we take the law of chemical mass action into consideration. Light acts not as a "stimulus" but acts by increasing the mass of certain chemical compounds, and it is the mass of these products which is responsible for the effect of light. Now, mass action is not proportional to the rapidity of the *change* of acting masses but to the acting mass itself. When two sides of an organism are struck by light of different intensity the quantity of photochemical products on both sides becomes unequal. In galvanotropism the galvanic current alters the distribution of the mass of certain ions along the nerve elements.

It can be shown that gravitation acts by influencing the distribution of chemical substances in an organism. When the stem of a plant is put into a horizontal position certain chemical substances gather in greater concentration on the lower side of the stem; and this causes a difference in the velocity of chemical reactions between the lower and the upper side. As a result of this we notice the bending. In the normal upright position of the plant the same substances were distributed equally about the axis of symmetry.

The following facts may be offered as a proof for this statement.⁵²⁶ When we put a piece of the stem of *Bryophyllum calycinum* in a horizontal position it soon bends and gradually assumes the form of a U with the concave side above (Fig. 35). This bending is due to the fact that the cortex on the under side of the stem grows in length while the cortex on the upper side remains unaltered

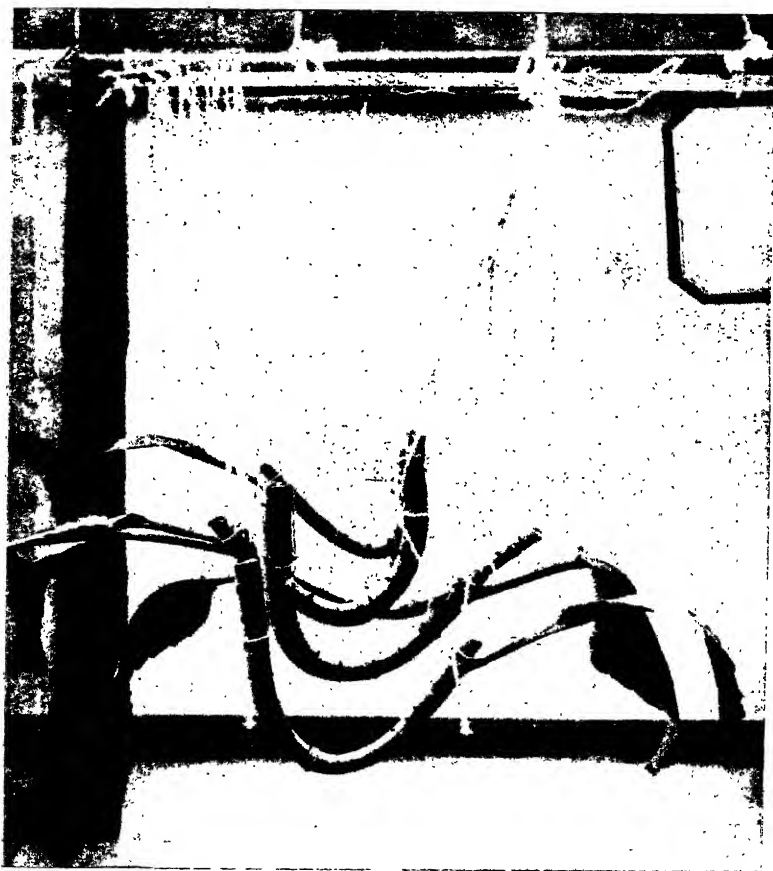


FIG. 35.—Geotropic curvature of stems of *Bryophyllum calycinum*. These stems were originally straight and suspended in a horizontal position. In about ten days they bent, becoming concave on the upper side. The black rings, made with india ink, which were originally parallel, remain unaltered on the upper side of the stems, while their distance increases on the lower side, indicating that the curvature is due to an increase in growth on the lower side (of the cortex) of the stem.

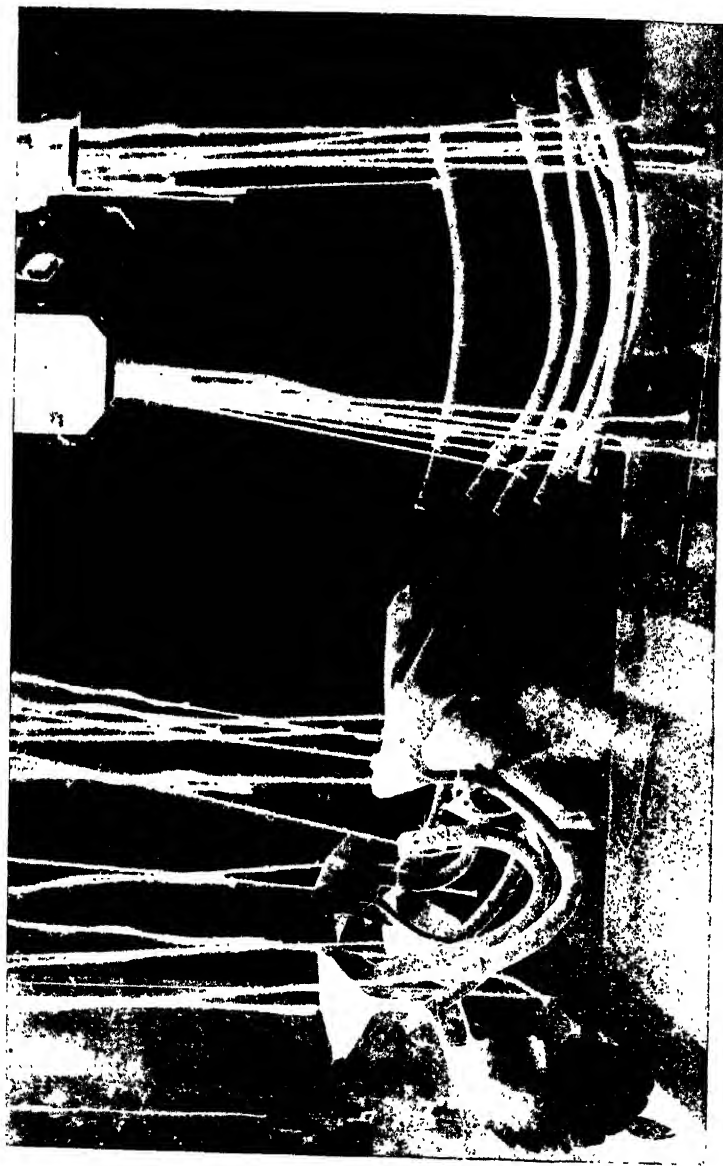


FIG. 36.—All stems were originally straight and suspended horizontally. The stems to the right, without leaves, bend much less in the same time than the stems to the left, with two apical leaves; the leaves supplying material for growth to the cortex of the stem on which the bending depends. This material collects in greater masses on the lower side of the cortex than on the upper side.



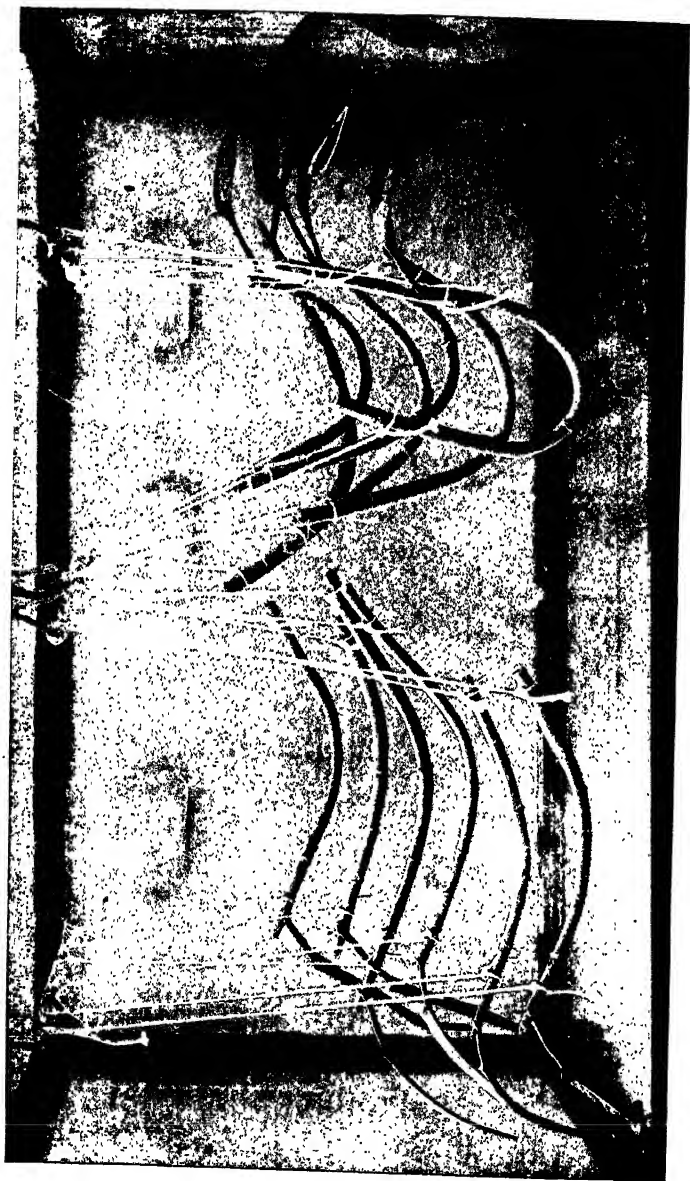


FIG. 38.—In stems to the left the size of the leaves is reduced by cutting off the lateral parts of the leaves. Stems to the right possess intact leaves. Notice that the geotropic curvature in the latter is much stronger than in the former.

(Loeb ⁵⁴²). This can be demonstrated if we mark the cortex in definite intervals with india ink at the beginning of the experiment (Fig. 35). After some time the distance between these marks will increase in a certain region of the under side, while it remains constant on the upper side, and this difference causes the bending. This positive increase in length of the under side can only happen through growth, and this growth of the cortex on the lower side of the stem takes place at the expense of material furnished constantly by the leaves which send it in the direction toward the basal end of the stem. When we compare the rate of geotropic bending of horizontal stems without leaves and with one or two leaves at the apex, we find that the bending in the latter is much more rapid (Fig. 36), owing to the greater mass of material supplied for the growth of the cortex, and the same is true, if we compare the rate of curvature of stems having a whole apical leaf attached with that of stems having an apical leaf whose mass has been reduced by cutting off parts of the leaf (Figs. 37 and 38). The writer has shown in other experiments that under equal conditions leaves produce material fit for growth in proportion to their mass. It is, therefore, a safe inference that the influence of the mass of an apical leaf upon the rate of geotropic bending is due to the mass of material it sends into the stem. This material has obviously a tendency to behave like a liquid—which it probably is—and to sink to the lower level. It is, therefore, useless to look for a “gravitational stimulus.” ^{526, 544}

What has been demonstrated in this case explains probably also why the apex of many plants when put into a horizontal position grows upward, and why certain roots under similar conditions grow downward. It disposes

also in all probability of the suggestion that the apex of a positively geotropic root has "brain functions." It is chemical mass action and not "brain functions" which are needed to produce the changes in growth underlying geotropic curvature.

2. As long as animals are in such a position that their plane of symmetry goes through the center of the earth, the position of their eyes and limbs is symmetrical in regard to their plane of symmetry. If, however, we incline the animal we can bring about forced movements and forced changes of position of the same nature as those caused by injury of one side of certain parts of the brain. Thus we have seen that if we cut the left side of the medulla oblongata in a shark, its two eyes are no longer in a symmetrical position but the left eye looks down and the right eye up, when the shark is kept in a normal position. The same change can be brought about in a normal shark by the influence of gravitation. When the shark is kept in a position with its right side inclined downward, the right eye is turned upward, the left eye downward. This has nothing to do with light or vision, since it occurs in the dark just as well as in an illuminated room. The abnormal position of the eyes lasts as long as the animal is kept in this abnormal position. The experiment shows that if the plane of symmetry is no longer vertical, forced positions of the eyes can be produced of the same nature as those produced by one-sided injury of certain parts of the brain.

Just as in the case of one-sided injury to the medulla oblongata the changes in the position of the eyes are accompanied by changes in the position of the pectoral fins, so also when we put a normal shark with one side downward or half downward.²⁸⁹ If the right side of such

a fish is down the right pectoral fin is turned more ventrally, the left fin is turned more dorsally. This means, the tension of the muscles causing the right fin to press down and the left fin to press up is increased. This is the mechanism by which the normal "equilibrium" or more correctly the normal geotropic orientation of the animal is maintained. If the animal should accidentally roll to one side in its normal movements, the tension of the muscles of the pectoral fins would automatically change in such a way as to restore the normal orientation of the animal, whereby the plane of symmetry becomes vertical again. This "maintenance of equilibrium" is therefore a case of automatic orientation by gravitation comparable to the automatic orientation by light.

Geotropic changes in the position of the eyes are not confined to fishes,³²⁰ they can be demonstrated in a rabbit and in crustaceans as well.

In vertebrates the reactions leading to the maintenance of equilibrium are apparently produced in the ear, since they disappear if the acoustic nerves are cut. Moreover, those parts of the brain whose injury brings about such changes in the position of the eye and the fins are parts of the receiving fibers from the acoustic nerve.²⁹⁰

It seems that some change in the pressure upon the endings of the auditory nerve is responsible for the effects. There are fine grains of CaCO_3 —the otoliths—in the ear of many species pressing on the underlying nerve endings. If we put the median plane of a fish at an angle of 45° with the vertical, the otoliths will no longer press down equally in both ears. The idea first suggested by Delage that it is the pressure of the otoliths upon the nerve endings which is responsible for these reactions receives some support by a well-known experiment by

Kreidl.²⁷⁰ A crustacean, *Palæmon*, loses its otoliths in the process of moulting and the animal curiously enough replaces them by picking up small grains of sand and putting them into its ears. Kreidl kept such crustaceans in jars free from sand but containing fine particles of iron which the crustaceans after moulting put into their ears. He expected that a magnet would now influence the animals as powerfully as gravitation, and this was the case. When, *e.g.*, he brought a magnet from above and the right near the animal the latter turned to the left and downward. The animal, therefore, behaved as if changes of pressure of the otolith upon the nerve endings determined its geotropic orientation.

The theory meets with two difficulties which, however, are not insuperable. First, removal of all the otoliths does not interfere with the normal orientation of the animal. This might find its explanation in the fact that the eyes act as a substitute. Delage had shown that if the otocysts are removed in crustaceans or cephalopods the animals lose their normal orientation more easily when they swim about excitedly than do normal animals. In order to show the effects clearly, however, it was necessary to blind the animals. Animals which were merely blinded but had their otocysts did not show these disturbances of equilibrium.¹¹⁹

The second difficulty is the fact that animals which possess naturally no otoliths are yet able to show such geotropic reactions, *e.g.*, certain crustaceans like *Gelastimus* and *Platyonichus*. We may assume that the pressure of liquids on the nerve endings may have a similar effect as the pressure of the otoliths.

The next question is, How does the pressure on a nerve ending bring about changes in the tension of muscles?

We suspect that this occurs through a change in mass action in the nerve endings, in analogy to our experiments on the influence of the mass of the leaf on the geotropic curvature of *Bryophyllum*, but experimental data are lacking.

3. We observe phenomena of geotropism in animals which have no ears, but this need not surprise us in view of the observations on geotropism in plants, and in hydroids (*Antennularia antennina*). The writer²⁸⁹ had found that a holothurian (*Cucumaria cucumis*) has a tendency to creep upward when put on a vertical object until it reaches the highest level, where it remains. When put on a vertical plate of glass or slate, these animals creep untiringly upward if only the plate is turned 180° around a horizontal axis as soon as they have reached the highest point. It could be shown that light and oxygen supply have nothing to do with the phenomenon. Jennings observed that *Paramæcia* always gather at the highest point of a vertical tube and that they assume this position by active ciliary motion. Lyon³²³ assumes that the body of *Paramæcia* contains substances of different density whose location is changed by changes in orientation of the organism to the center of the earth and that these changes automatically turn the animal again so that its oral pole is directed upward. It will then continue to swim in this direction.

4. It is known since Knight's experiments that centrifugal force can act like gravitation and we must assume that the centrifugal force leads to an alteration in the distribution of the sap or of other substances in the cell. This leads to differences in the rate of chemical reactions and may account for the phenomena of orientation under the influence of centrifugal force.

When an animal, *e.g.*, a shark or a pigeon, is rotated on a turntable, *during* rotation a nystagmus is observed in the motions of the eyes and sometimes also of the head. If the rotation is not too rapid the eyes move slowly in the same plane but in an opposite direction from the rotation of the turntable, until they form a maximum angle with their normal position in the head; then they rapidly swing back and the whole phenomenon is repeated. This phenomenon is called nystagmus. It depends upon the nerve endings in the semicircular canals, but is not dependent upon the motion or pressure of the lymph in the canals,^{290, 319, 320} since the cutting out of the canals in the shark or the plugging up of the canals in the pigeon¹⁴¹ leaves the phenomenon unaltered. When after some rotation the motion of the turntable suddenly stops, a nystagmus of the eyes or head in the same plane but in the opposite direction as during the rotation is observed.

Maxwell⁵⁵⁴ has shown that if *Phrynosoma* is rotated on a horizontal plane with constant velocity and the eyes of the animal are closed, compensatory motions of the head are produced as soon as the angular velocity exceeds a certain value which was 8 seconds for a rotation through an angle of 45°.

CHAPTER XIV

FORCED MOVEMENTS CAUSED BY MOVING RETINA IMAGES: RHEOTROPISM: ANEMOTROPISM

THE experiments on forced movements show that we have three groups of forced movements, (1) right to left and left to right (circus movements); (2) forward movement, and (3) backward movement. The latter is not always possible. A fourth group, the rolling motions around the longitudinal axis may be omitted here in order to simplify the discussion.

The forced movements, called forth by the galvanic current, supported the idea that the nervous elements determining these motions must have a definite orientation and that this orientation bears some simple relation to the direction of motion caused by their activity. The experiments on the effect of blackening different parts of the eye indicate that the different parts of the retina of positively heliotropic insects are connected in a simple way with the main centers of the three types of forced movements: namely, the left eye is connected with the brain center causing motions from right to left (and the right eye with the center for the opposite motion); the lower halves of the retina with the forward movements, the upper halves with the backward movements.

We know through the work of Ewald Hering that each illuminated element on the human retina determines a definite motion of the two eyes which move as if they were a single organ, and that this motion is a function of the

location of the illuminated element in the retina. This fact induced the writer to suggest in his first publication on tropisms that the act of focussing in our vision was simply a phenomenon of heliotropism. "The general principle of orientation of organisms to light is also manifested in our act of binocular vision which results automatically in such an orientation of the two retinae that the image of the luminous point falls upon the two foveae centrales of the retinae" (which are symmetrical elements). In other words, when an object causes us to turn our eyes to it we are dealing with a phenomenon of forced (heliotropic) movement. In order to prove this it is necessary to show that a moving retina image can produce forced movements determined by the direction of motion of the luminous object. The difficulties inherent in the proof for such a statement lie in the general prejudice that the motions of an animal are directed to a purpose and it is, therefore, necessary to devise experiments which exclude the assumption of an interest on the part of the animal in the motion.

The writer observed years ago that when a fly is put on a rotating disk it rotates in the opposite direction from the disk. When the motion of the turntable ceases these compensatory motions of the fly stop also and none of the after effects mentioned at the end of the previous chapter are noticed.²⁸⁶ This suggested that the so-called compensatory motions of insects on the turntable have a different origin from that of vertebrates. The phenomenon was explained by Rádl, who proved that the compensatory motions of insects on the turntable are produced in the eye and that they are due to the fact that the eye tries automatically to fix the same object.⁴⁴⁷ This agrees with the observation of Lyon who had already

demonstrated previously that the compensatory motions of insects on a turntable stop when their eyes are blackened.³¹⁹ Such forced motions, due to the influence of the motion of the retina image, can be demonstrated in the Californian lizard *Phrynosoma blainvilli*, which is an ideal object for such experiments²⁹⁸ and in this animal it is possible to separate these effects from the compensatory motions caused by centrifugal force.

It was accidentally observed by the writer that when the lizard *Phrynosoma* is kept at the window of a moving train with its eyes toward the window, a nystagmus of the head of the lizard ensues, the head moving slowly in a direction opposite to that of the moving train, as if to keep its eyes fixed on the objects outside—telegraph poles and trees, etc. The head moves until it is bent maximally, when it is brought back into its normal position with a quick jerky movement, and then follows again the apparent motion of the objects outside, and so on. These nystactic motions last for hours, in fact as long as the animal is kept with its head toward the window. As soon as it is turned around so that it cannot see the objects outside, the nystactic motions of the head cease. When the animal is put on a turntable and rotated slowly, vigorous compensatory movements can also be observed during rotation. If, however, the eyes of the lizard are closed during rotation these movements are considerably diminished though they do not cease entirely. They are also considerably diminished when the animal with its eyes open is rotated on a turntable surrounded by a high gray cylinder of cardboard which excludes the possibility of images of outside objects moving on the retina. We can also produce compensatory motions of the head if the animal

is kept quiet and objects are moved in front of it, the eyes following the moving object.

It is of interest to separate the nystagmus or compensatory motions of eyes and head caused by the orienting effect of a moving retina image from those caused by the orienting effect of centrifugal force and this can be done easily in *Phrynosoma*.

When the lizard is rotated very *slowly* on a turntable with *its eyes closed*, only very slight compensatory motions of the head and body are observed during rotation, while very powerful compensatory motions are produced when the motion of the turntable is suddenly interrupted after a rotation lasting about thirty seconds.

When, however, the same experiment is made with *the eyes of the lizard open* the reverse is observed. The compensatory motions of the animal during rotation are exceedingly vigorous, while the compensatory motions of the animal after the interruption of the rotation are slight.

When the eyes of the animal are closed we are dealing only with the geotropic effect of passive rotation; when the eyes are open the orienting influence of the moving retina image is added algebraically to the orienting effect of centrifugal force upon the ear. These two influences act in the same sense *during* rotation and therefore are additive; while *after* the rotation they act in the opposite sense to each other. When we rotate the body of an animal passively to the right, *during* rotation the objects have an apparent motion to the left and the eyes and head of the animal are compelled to follow these moving objects, *i.e.*, to the left. The geotropic effect of passive rotation of the animal to the right also causes a motion of the eyes and head to the left and hence both effects are additive.

When a human being has been rotated passively to the right for some time, at the interruption of the passive motion the eyes move slowly to the right and return rapidly to the left. Only the slow motions give rise to the sensation of an apparent motion of the objects and hence after the sudden stopping of a passive rotation to the right the objects seem to such a person to move to the left. The geotropic after effect, after passive rotation to the right, consists in inducing passive compensatory motions to the right, *i.e.*, in the opposite sense of the orientation caused by the apparent motion of the visual objects. Hence in the after effect the orienting effect of the retina image and the centrifugal effect weaken each other.

Lyon ^{321, 322, 326} has shown that the phenomena which were formerly described as rheotropism in fish are due to the orienting effect of moving retina images. The reader is familiar with the fact that many fish when in a lively current have a tendency to swim against the current. This phenomenon was believed to be due to the friction of the water. Lyon showed that fish orient themselves just as well when they are put into a closed glass bottle, which is dragged through the water, although in this case they are not under the influence of any friction from the current. When the bottle is not moved the fish swim in any direction inside the bottle. It is obviously the motion of the retina images of the objects on the bank of the brook which causes the "rheotropic" orientation of fish. When driven backward by the current or when dragged backward in a bottle through the water, the objects on the bank of the river seem to move in the opposite direction. The animal being compelled to keep

the same object fixed, an apparent forward motion of the fixed object changes the muscles of the fins in such a sense as to cause the animal to follow the fixed object automatically.

When such rheotropic fishes were kept in an aquarium and a white sheet of paper with black stripes was moved constantly in front of the aquarium the fish oriented them-

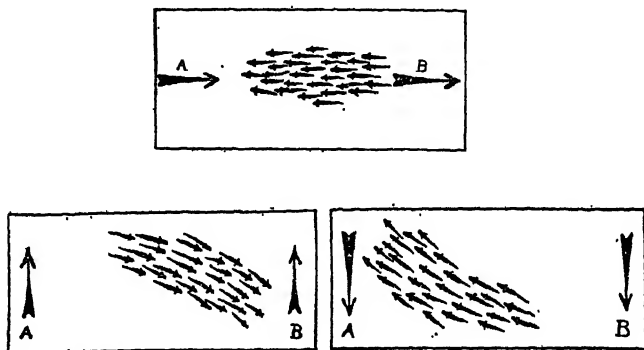


FIG. 39.—Influence of motion of the hand of an observer on the direction of the motion of a swarm of sticklebacks in an aquarium. The arrows indicate the direction in which the hand was moved. The swarm of fish moves always in the opposite direction in which the hand is moved. (After Garrey.)

selves against the direction in which the paper and its stripes moved. The phenomenon was more marked in young than in older specimens.

All the phenomena of rheotropism ceased in the dark or when the fish were blind.

Wheeler⁵⁰⁸ has observed a phenomenon of anemotropism, namely that certain insects have a tendency to put the axis of their body in the direction of and against the wind. He considers this analogous to the phenomenon of rheotropism in fishes. The cause is also in all probability the tendency toward fixation of the moving retina image.

A very pretty demonstration of the orienting effect of moving retina images was discovered by Garrey in

sticklebacks.¹⁷⁶ When a swarm of such fish was kept in an aquarium it was noticed that all the fish were oriented with the long axes parallel and that the whole school swam in a course parallel, but in a direction opposite, to that of the moving observer. If the observer remains stationary opposite the aquarium and moves an object, preferably white, which is held in the hand, the little fish at once respond by moving slowly and oppositely to that of the moving object. They can be thus made to move up or down or to the right or left (Fig. 39).

By experiments which space forbids us to report in detail Garrey has reached the conclusion that the motion of a near object causes an apparent motion of the whole horizon in the opposite direction and this apparent motion the fish tries to compensate by the motions of its body. This brings the observations on the stickleback into harmony with the general influence of moving retina images, consisting in a compensatory motion of the fish.

We have already referred to the fact that the influence of a moving retina image is capable of compensating the forced movement of a dog after a one-sided lesion of the cerebral hemispheres.

CHAPTER XV

STEREOTROPISM

Our orientation in space is determined by three groups of tropistic influences, two of which we have already discussed, light and gravitation. The third one is pressure on certain nerve endings of the skin. When the tactile influences on the skin of the soles of the feet are weakened (as is the case in locomotor ataxia), the patient finds it difficult to stand and walk in the dark. When he can use his eyes the difficulty is diminished, since the orienting effect of the retina image can compensate the tactile deficiency; just as we have seen that the effect of the loss of the ears in crustaceans can be compensated by the orienting influence of the eyes.

The rôle of tactile influences on the orientation of animals is most clearly demonstrable in starfish, flatworms, and many other animals, when put on their backs. The animals "right" themselves, *i.e.*, they turn around until the ventral surfaces or their feet are pressed against solid objects again. As the writer pointed out long ago,²⁹³ gravitation has nothing to do with the phenomenon, since starfish will stick to solid surfaces with their tube feet even if by so doing their backs are permanently turned to the center of the earth. Unless the nerve endings on the sole of their tube feet are pressed against a solid surface the animals are restless and the arms move about until the feet are again in contact with solid bodies. This phenomenon of orientation the writer called stereotropism.

Quantitative investigations of this form of tropism are

still lacking and we must be satisfied with a few descriptive remarks.

Certain animals show a tendency to bring their body completely into contact with solid bodies, *e.g.*, by creeping into crevices. Without further experimental test this might appear as an expression of negative heliotropism, but it can be shown that this assumption would be wrong. *Amphipyra* is a positively heliotropic butterfly which, in spite of its positive heliotropism, shows the peculiarity that it creeps into crevices when given an opportunity. Such animals were kept in a box at the bottom of which was a square glass plate resting with its four corners on supports just high enough to allow the animals to creep under the glass plate. After some time every *Amphipyra* was found under the glass plate. This happened also when the glass plate was exposed to full sunshine, while the rest of the box was in the shade.²⁸⁷

The same stereotropism is found in female ants at the time of sexual maturity. When such animals are put into a box containing folded pieces of paper or of cloth, after some time every individual is found inside the folds. This happens also when the boxes are kept in the dark.²⁸⁷

The same form of stereotropism is found in many species of worms. When earthworms are kept in jars with vertical walls they are found creeping in the corners where their body is as much as possible in contact with solid bodies. It is this tropism which compels the animals to burrow into the ground.

Maxwell³⁴⁹ kept *Nereis*, a form of marine worms, which burrows in sand, in a porcelain dish free from sand. Into the dish glass tubes were put, whose diameter was of the order of that of the worms. After 24 hours every tube was inhabited by a worm who made it its permanent

abode. They even remained in the tube when exposed to sunlight which rapidly killed them.

We find the opposite, negative stereotropism, in many pelagic animals, *e.g.*, larvæ of the barnacle or of other crustaceans, which avoid contact with solids. The phenomenon is liable to interfere with heliotropic experiments.

The importance of stereotropism in animals was first pointed out by the experiments of Dewitz on the spermatozoa of the cockroach.^{120, 121} He noticed that when a drop of salt solution containing the spermatozoa was put under a cover glass resting on low supports on a slide, the spermatozoa collect at the solid surfaces of the slide and cover glass, while the liquid between remains free from spermatozoa. When a small glass bead is put into the liquid the spermatozoa will also swim on the surface of the bead, never leaving it again. Dewitz is of the opinion that this stereotropism is of assistance in securing the entrance of a spermatozoon into the egg. The egg of the cockroach is rather large and the spermatozoon can enter it only through a micropyle. When the egg is laid it passes by the duct of the seminal pouch in which the female keeps the sperm after copulation. On passing the duct some spermatozoa reach the egg. Dewitz points out that these cannot leave the surface of the egg any more but are compelled to move incessantly on the surface of the egg until one of the spermatozoa by chance gets into the micropyle.

It is an important fact that different organs of the same organism react differently. We have already mentioned the tendency of starfish or flatworms to right themselves, *i.e.*, their ventral surface is positively their dorsal negatively stereotropic. The stolons of hydroids stick

to solid bodies, while the polyps bend and continue to grow away at right angles from solid bodies with which they come in contact. Thus the stem of *Tubularia mesembryanthemum*, a marine hydroid, grows in a straight line. When such stems, after their polyp is cut off, are put with one end in sand, the free end forms a new polyp and the stem continues to grow in a vertical direction upward. When, however, the stem is put near the glass wall as soon as the polyp grows out it bends away from

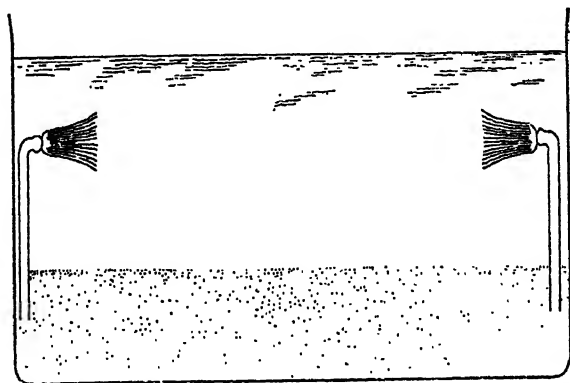


FIG. 40.—The regenerating polyp of *Tubularia* when in contact with the glass wall of an aquarium bends at right angles to the glass wall.

the solid wall, and the stem will now continue to grow at right angles to the vertical wall (Fig. 40).

This phenomenon raises the question whether or not the law of chemical mass action underlies phenomena of stereotropism. We have seen that this law dominates the phenomena of heliotropism, inasmuch as the Bunsen-Roscoe law is the expression of the influence of light on the mass of the photochemical reaction product. We have also been able to show that in the case of the geotropic curvature of *Bryophyllum* the mass of the apical

leaf determines the rate of geotropical curvature of a horizontally placed stem. The only way in which the mass of the leaf could have such an influence is through the mass of substances it sends into the stem, so that this case of geotropism is a function of mass action. There are indications that the way contact with a solid influences the behavior of living matter is also through the influence on the rate of certain chemical reactions. The writer observed that the stolons of a hydroid, *Aglao-phenia*, have a tendency to adhere to solid surfaces and not to leave them any more if they once reach them, and that as soon as such a stolon reaches a solid surface, *e.g.*, a piece of a glass slide, its growth is accelerated considerably. It was very astonishing to notice how much more rapid the growth of roots of *Aglao-phenia* was when they were in contact with a solid body than when they grew in sea water. The rate of growth is the function of a chemical mass action (Loeb⁵⁴³).

CHAPTER XVI

CHEMOTROPISM

1. When we create a center of diffusion in water or in air we may theoretically expect orienting effects. Thus when a fine capillary tube containing a solution of a salt, *e.g.*, sodium malate, is put into a drop of water containing motile organisms, and the right side of an organism is turned to the source of diffusion, the diffusing molecules will collect in increasing concentration on that side. On the left side of the organism, no such increase in the concentration of molecules will occur. If now the molecules collecting on the right of the organism in increasing density are able to produce some chemical or some concentration chain effect, the two sides of the organism will be acted upon unequally and the tension of the symmetrical motile organs will no longer be the same. As a consequence the organism will turn until the mass of molecules or ions striking the organism in the unit of time will be the same for both sides. These effects only take place when the organism is close to the opening of the capillary tube, since the diffusion from the tube is slow.

It is obvious, however, that it is difficult to provide experimental conditions which give exact chemotropic reactions. First of all, if the diffusion is rapid the differences in concentration of the effective chemotropic substance on two sides of an organism are too slight to result in a turning movement. A second condition which is liable to vitiate the result are the unavoidable convection currents due to changes or differences of temperature. In

order to get clear results a method must be used which prevents a rapid diffusion of the substance; and, moreover, the current of diffusion must be confined to an almost straight line. It is possible that Pfeffer's method satisfies this condition.^{424, 425} He introduced the substance to be tested for its chemotropic effect into a capillary tube, the end of which was then sealed. The other end was pushed into a drop of water containing the suspension of the organisms whose chemotropism was under investigation. From this capillary the diffusion was extremely slow. Moreover, the current of diffusion was approximately linear at the orifice. Hence the test for the existence of positive chemotropism was perhaps possible. When an organism, struck sidewise by the line of diffusion near the opening of the capillary tube, turns toward the tube going into it, some probability of positive chemotropism exists; and when all the organisms coming near the orifice of the tube are thus compelled to go into it, the probability may become certainty, provided that the substance used does not paralyze the organism and therefore act as a trap, allowing the organisms to come in but not to go out. The capillary tubes used were of 10 to 15 mm. length and of a width of about 0.1 mm. Pfeffer and his pupils found that the spermatozoa of ferns go in large numbers into a capillary tube containing sodium malate in a concentration of 0.01 per cent. (a solution ten times as diluted is still slightly active). This effect of the malate is specific in this case and this indicates that either a definite chemical action of the malate ion or a specific permeability of the organism for it is the source of the chemotropism. Such specific chemotropic effects are not rare, since Pfeffer found that *Bacterium termo* and *Spirillum undula* are positively chemotropic

to a liquid containing 0.001 per cent. of peptone or of meat extract. It is stated that cholera bacilli are strongly attracted by potato sap. Pfeffer found also that the spermatozoa of certain mosses are positively chemotropic to cane sugar solution in dilutions of 0.1 per cent.

Pfeffer's work preceded the discovery of electrolytic dissociation, and his pupils Buller⁸⁹ and Shibata⁴⁶⁵ made some of the additions required by the theory, namely, that it is the malate anion which acts in the case of the spermatozoa of the ferns, and that when the anion is offered in the form of malic acid the H ion counteracts the effect of the malate anion.

Shibata made extensive experiments on the chemotropism of the spermatozoa of *Isoëtes*⁴⁶⁵ which he found positively chemotropic for the malate anion, and also for the succinate, tartrate, and fumarate anion, when offered in the form of their neutral salts. The anion of the stereoisomere of fumaric acid, namely of malëic acid, was without effect. This indicates a high degree of specificity of these reactions. Neutral sodium malate acted best in dilutions from m/100 to m/1000, but some action could still be discovered in m/20,000 solutions.

When malic acid was used no positive chemotropism could be discovered in solutions of m/100 or above on account of the contrary effect of the hydrogen ion, and the spermatozoa of *Isoëtes* did not even go into capillary tubes containing m/1000 malic acid. When any acid other than malic was added to sodium malate the motion of the spermatozoa into the tube was prevented, even a m/6000 HCl solution still had such an effect.

Shibata studied especially the mode by which the spermatozoa are oriented chemotropically by malates and found that the reaction consists always in a turning of

the axis of the body of the spermatozoa toward the capillary tube containing malates or succinates, as the tropism theory demands.

When the capillary tube and the surrounding medium contain the same solute for which the organisms are positively chemotropic, they will not go into the tube unless the concentration in the tube is a *definite* multiple of the concentration of the outside solution. Thus Pfeffer found that the concentration of sodium malate in the capillary must be at least thirty times as great as in the outside solution to induce the spermatozoa of fern to move into it, and in the case of *Bacterium termo* the solution of meat extract in the tube had to be at least four times as great as the outside solution. In the case of *Isoëtes* spermatozoa Shibata found the ratio of about 400 to 1. This constancy of the ratio is known as Weber's law, which therefore holds for chemotropic phenomena.

Lidforss²⁸¹ found with the aid of Pfeffer's method that the spermatozoa of *Marchantia* are positively chemotropic to certain proteins, especially egg albumin, vitellin from the egg yolk, hemoglobin, and mucin of the submaxillary gland; blood albumin, casein, and legumin were less effective. The lowest concentration for hemoglobin solutions and for egg albumin was 0.001 per cent.!

It may also be stated that Lidforss found a chemotropic effect of proteins upon the direction of growth of pollen tubes.²⁸⁰

Bruchmann⁸¹ found that the spermatozoa of *Lycopodium* were positively chemotropic to the watery extract in which pieces of the prothallium had been boiled. Pfeffer's capillary method was used. They showed also positive chemotropism to the citrate anion. Thus, sodium citrate was efficient in a 0.1 to 0.5 per cent. solution. The

lower limit was a little above a 0.001 per cent. solution. The effect of the free citric acid was a mixed one since the spermatozoa were negative to H ions and positive to the citrate anion. Instead of being able to use a 0.1 per cent. solution, as in the case of the sodium salt, a 0.01 per cent. solution was the highest concentration to which they were positively chemotropic. This means that the hydrogen ion of citric acid solutions above $m/1000$ repel the spermatozoa, while when solutions of $m/2000$ or below are used the hydrogen ion effect no longer inhibits the positive effect of the citrate anion. In addition the validity of Weber's law could be demonstrated. The spermatozoa were indifferent to malates, oxalates, and many other salts, as well as to sugar and proteins.

2. While all the botanical observers, from Buller on, had found that the hydrogen ion has only a preventive effect upon the positive chemotropism of lower organisms, Jennings tried to show that acids have a positive effect, especially when in low concentrations.²⁵⁰ But his concentrations are not quite as low as he seems to assume, since a 1/50 per cent. ($m/180$) HCl solution, toward which he believes to have proven positive chemotropism of *Paramæcia*, is a deadly concentration.^a Jennings's interest in the problem was aroused by a phenomenon of aggregation, not infrequently found in the suspensions of infusorians.

It is well known that when certain infusoria are left undisturbed they do not remain scattered, but gather in more or less dense groups. Thus, if they are mounted on a slide in a thin layer of water, soon dense aggregations will be formed in certain areas, while the remainder of the

^a The cells of the stomach resist a much higher concentration of HCl but this is an exception. Infusorians, fish, and organisms in general are killed in a short time in $m/180$ HCl or in a much lower concentration of acid. Thus *Fundulus* does not live more than one hour in $m/3000$ HCl or HNO_3 . (Loeb, J., and Wasteneys, H., *Biochem Z.*, 1911. xxxiii, 489; 1912, xxxix, 167.)

slide will be nearly deserted. One of the first investigators to describe this phenomenon was Pfeffer. He observed its occurrence in *Glaucoma scintillans*, and less markedly in *Colpidium colpoda*, *Stylonychia mytilus*, and *Paramecium*. Pfeffer was inclined to believe that these aggregations were due, partly at least, to a contact stimulus, resulting from a striking of the organisms against small solid bodies, and especially against each other.²⁵⁰

This conclusion of Pfeffer may after all be correct, since it has been shown that sea water containing jelly from the egg of a sea urchin causes spermatozoa to stick together for some time when they impinge upon each other. This agglutination no longer occurs when the spermatozoa are immobilized. Jennings came to the conclusion that these aggregations of infusorians are due to the fact that they can go into a weak concentration of acid, while they cannot escape from such a weak concentration; and since *Paramecia* themselves produce CO₂, he assumed that the CO₂ produced by themselves acts as a center of attraction for other *Paramecia*. In order to prove this he used the following method:

The organisms were studied in a thin layer of water, by mounting them on a slide covered with a large cover glass supported near its ends by slender glass rods. Their reactions were tested by introducing with a capillary pipette a drop of the substance in question beneath the cover glass, or in some cases by allowing it to diffuse inward from the side of the cover glass.²⁵⁰

Thus Jennings introduced a drop of 1/50 per cent. (m/180) HCl on a slide containing *Chilomonas*. Very soon a somewhat denser ring of these individuals was formed around the drop (Fig. 41). A 1/50 per cent. HCl solution paralyzes (and soon kills) *Chilomonas* or *Paramecia* and hence the surface of the drop must act like a trap into which the organisms will steadily swim, without being able to swim back. This will naturally increase

the density of organisms around the drop and may give rise to a ring formation around a high concentration of HCl although the organisms are not positive to the acid. Jennings found, however, that when such organisms are in a drop of weak acids which do not paralyze the organisms quickly, *e.g.*, 1/50 per cent. acetic or in CO₂ solutions, they become negative to the surrounding neutral medium (H₂O or hay infusion) and stay in the acid. He, therefore, assumes that the organisms are positive to weak acid, and

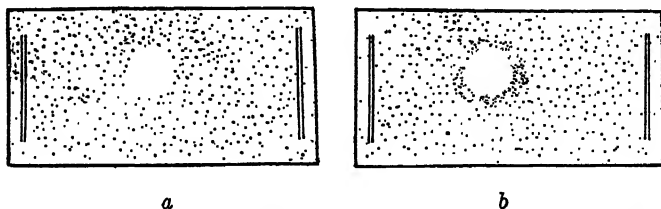


FIG. 41.—Reaction of *Chilomonas* to a drop of 1/50 per cent. HCl. *a*, preparation immediately after the introduction of the drop (no organisms either within or gathered about the drop). *b*, the same preparation a few minutes later. (After Jennings.)

negative to strong acid as well as to their natural neutral or faintly alkaline medium.

This negativity to their natural surroundings when in weak acid as well as to strong acid when in weak acid Jennings does not interpret in terms of the tropism theory, and in this he is probably correct. He interprets both phenomena as a trap action due to the asymmetry of certain infusorians; a sudden change in the concentration of a solution causes a reverse of the stroke of their cilia by which the organism is driven back. When the old normal stroke of the cilia is resumed the direction of the locomotion is changed on account of the asymmetrical arrangement of the cilia. This happens when the organisms go from weak into strong acid or from weak acid into

a neutral medium. In this way a collection of the organisms at the surface of a drop of acid may be brought about. This phenomenon is not tropistic in the strict sense of the word, and as a matter of fact *Paramæcium* is not positively chemotropic to acid of any strength.

Barratt^{2a} investigated the chemotropism of *Paramæcia* for varying concentrations of different acids with

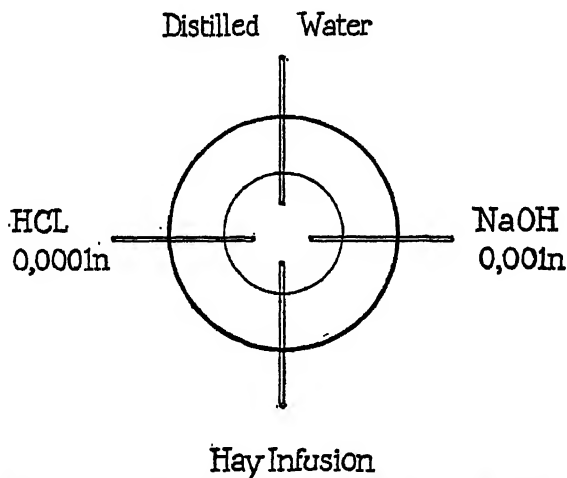


FIG. 42.—Method of proving that *Paramæcia* are not positive to acids of low concentration. (After Barratt.)

Pfeffer's method of capillary tubes, counting the number of individuals going into the tube containing acid and comparing it with the number going simultaneously into a control tube containing only distilled water free from CO_2 (Fig. 42).^b The acids used varied from 0.001 N to 0.0001 N. The results were unequivocal. Toward solutions of 0.001 N the *Paramæcia* are negative and possibly

^b In addition two other controls accompanied the test, namely, one tube containing hay infusion (the natural medium of the organisms) and one alkali.

also slightly negative to acids as weak as 0.0001 N. In no case, not even with the weakest acid, was it possible to prove the existence of positive chemotropism for acid (or base). The number of *Paramæcia* which went into a tube containing, e.g., 0.00002 N acid, was on the average not greater than that which went into the control tubes. The tubes were sufficiently wide so that the *Paramæcia* could and did move into the tubes. Barratt, therefore, concludes that acids have only a repelling action upon *Paramæcia* which, however, diminishes or disappears when the hydrogen ion concentration approaches that of distilled water.

The observations of Barratt contradict the statement that *Paramæcia* are positive to weak acid. We have seen that when spermatozoa or swarmspores are positive to malates this can be elegantly shown by Barratt's method. The same method has shown that when even a trace of acid is added to the neutral malates this positivity disappears. By testing systematically all concentrations of different acids within the range to be considered, Barratt found no trace of any positivity to or any trap action by weak acid for *Paramæcia*. It may be true, however, that when the organisms are in very dilute acid neutral or faintly alkaline water repels them in the way described by Jennings.

Barratt states also that there is nothing to support Jennings's assertion that the CO_2 given off by the *Paramæcia* causes the aggregation in their natural medium, since they are not positive to low concentrations of hydrogen ions. The natural aggregations of infusorians may be due, as Pfeffer suggested, to transitory agglutinations when *Paramæcia* impinge upon each other, and the stickiness or tendency to agglutinate may possibly be increased

by certain substances produced and excreted by the organisms themselves, *e.g.*, CO₂.

3. The results obtained with the spermatozoa of ferns and mosses by Pfeffer and other botanists led some authors to the tacit assumption that the spermatozoa of animals were positively chemotropic toward substances contained in or secreted by the eggs of the same species. Some accepted this assumption without test, others made tests which they considered adequate but which seem doubtful, and it may be of some interest to discuss the subject, since far-reaching conclusions might be based on these experiments. Pfeffer's method of testing for chemotropism with the aid of the capillary tube has proved satisfactory and the application of this method has shown that the spermatozoa of certain animals, *e.g.*, of sea urchins, are not chemotropic toward substances contained in or given off by the egg. Thus Buller, who had worked in Pfeffer's laboratory on the chemotropism of the spermatozoa of ferns, investigated carefully and extensively the question whether or not the spermatozoa of echinoderms are positively chemotropic for egg substances.⁹⁰ His results were entirely negative. Thoroughly washed, ripe unfertilized eggs of *Arbacia* (Naples) were put into a small volume of sea water for from 2 to 12 hours.

Capillary glass tubes, about 12 mm. long and 0.1 to 0.3 mm. internal diameter, and closed at one end, were then half filled with the (supernatant) sea water (which had contained the eggs) by means of an air pump. The tubes were then introduced into a large open drop of sea water, in which fresh, highly motile spermatozoa were swimming. If the eggs excrete an attracting substance it was argued that it should be present in the tubes, and the spermatozoa should collect there. . . . No attraction into the tube could be observed. Except for a surface-contact phenomenon to be further discussed, they went in and out with indifference. Apparently, therefore, the water which had contained the eggs exercised no directive stimulus on the spermatozoa whatever.

I then attempted to find some substance which could give a chemotactic stimulus to spermatozoa. The substances tested were such as are known to give a directive chemical stimulus to many protozoa, the spermatozoa of ferns, pollen-tubes, etc. The following solutions were tried by the capillary tube method: distilled water; meat extract 1 per cent.; KNO_3 10 per cent., 2 per cent.; NaCl 5.8, 2.9, 0.58 per cent.; K_2 malate 1, 0.1 per cent.; asparagin 1 per cent.; glycerine 5 per cent.; grape sugar 18, 9, 4.5, 2.25 per cent.; peptone 1 per cent.; alcohol 50, 25, 10 per cent.; diastase 1 per cent.; oxalic acid 0.9, 0.09, 0.009 per cent.; nitric acid 1, 0.1, 0.01 per cent.

No definite chemotactic reaction—neither attraction nor repulsion—was observed in any case. Into tubes containing the weaker solutions the spermatozoa went in and out with apparent indifference. . . . On coming into contact with strong acid solutions (oxalic acid 0.9, 0.09 per cent.; nitric acid 1, 0.1 per cent.) the spermatozoa *were killed, and thus formed slight collections*. They were thus not able to avoid acids by means of a negative chemotactic reaction.⁹⁰

Other authors, *e.g.*, Dewitz and the writer, have also reached the conclusion that the egg of the sea urchin contains no substance for which the spermatozoon of the same species is positively chemotropic, and that Buller's conclusions that positive chemotropism plays no rôle in the entrance of the spermatozoon of sea urchins into the egg is correct.

F. Lillie has recently expressed the opposite view, namely that the egg of the sea urchin contains a substance to which the spermatozoa are positively chemotropic and to which he gave the name "fertilizin."²⁸³ He first tried Pfeffer's correct method with capillary tubes with negative result, just as Buller and the rest of the observers. Instead of concluding that the spermatozoa are not chemotropic he discarded the method and used Jennings's method, stating that it gives "incomparably more delicate results than Pfeffer's method of using capillary tubes" (p. 533). Lillie found with this method that the spermatozoa of *Arbacia* are positively chemotropic to

H₂SO₄ of a concentration as high as N/10 and that they are never negatively chemotropic, not even to the highest concentrations of the strongest acid. It seems to the writer that Lillie's observations are more naturally explained on the assumption that when an acid is sufficiently strong and concentrated, *e.g.*, N/10 HNO₃ or H₂SO₄, it will paralyze and kill the spermatozoa, and that when a drop of such acid is introduced in sea water containing spermatozoa, a somewhat denser ring of the organisms will be formed around the surface of the drop on account of this action of the acid.

With the same method Lillie tried to prove that the spermatozoa of *Nereis* and *Arbacia* are positively chemotropic to extracts of their own eggs.²⁸³ He proceeded as follows: A suspension of *Arbacia* sperm, freshly made, was put under a raised cover slip and a drop of the supernatant sea water which had been standing over eggs (as in Buller's experiments) was introduced under the cover slip. Observation with the naked eye showed that around this drop of egg-sea water immediately a dense ring of spermatozoa formed and behind this a clear external zone was formed about 1.2 to 2 mm. wide. The dense ring then broke up into small agglutinated masses. In Lillie's opinion the formation of this dense ring of spermatozoa at the periphery of the egg-sea water is the expression of a positive chemotropism of the spermatozoa for a substance contained in the egg-sea water, the "fertilizin." He assumes that the spermatozoa near the drop of egg-sea water all swim to the egg-sea water, leaving a clear space behind them. While this explanation of the ring formation might be true—if supported by a direct chemotropic method like Pfeffer's—it can be shown that the ring formation is in all probability due to an entirely different

phenomenon which has no relation to chemotropism or any other tropism.

Buller had already observed that the supernatant sea water of sea urchins contains a substance which causes the agglutination of spermatozoa.⁹⁰

A drop of sea water in which eggs had been deposited was placed upon a slide and a drop containing spermatozoa near it. On joining the drops a large number of small balls were formed in a very few seconds. When very numerous spermatozoa were present the balls became 0.1 mm. in diameter, containing many thousands of spermatozoa packed together in a dense mass.

Buller explains the phenomenon as being due to small bits of egg jelly floating in the sea water

so small that they will (like spermatozoa) pass through ordinary filter paper and, so transparent that one cannot directly see them. A few spermatozoa become attached to each piece of jelly, the presence of which may be inferred from the manner in which the small groups of spermatozoa move about. Owing to the length of the spermatozoon, although its head may be imbedded in a jelly particle, the tail may remain partly free. The little collections of spermatozoa thus move about hither and thither in no particular direction. When two such groups come by accident into contact they fuse. Certain of the spermatozoa adhere to both little masses of jelly and lock them together. The fused mass combines with other simple and fused masses, and so on.^c

The writer was able to show that when the jelly of the egg of *Strongylocentrotus purpuratus* is dissolved by an acid treatment the eggs when washed and transferred to sea water no longer give off agglutinating substances, while the acid sea water containing the dissolved jelly, when rendered neutral through the addition of alkali, will cause the agglutination of sperm.³⁰² While all the jelly can be washed off with an acid treatment in the egg of *purpuratus*, the same is not true for the egg of *Arbacia*

^c This explanation of the fusion of two clusters to a larger one is perhaps not correct. The writer is inclined to ascribe it to the adhesion or agglutination of the spermatozoa of two neighboring clusters with each other, due to a sticky surface on the sperm head.

of Woods Hole. Here the acid treatment does not as a rule dissolve all the jelly, or possibly some new jelly may be given off by the egg.

While Buller may be correct in assuming that microscopic pieces of the egg jelly form the center of these sperm clusters, the writer reached the conclusion that the dissolved mass of the jelly makes the surface of the spermatozoa transitorily sticky, so that if they impinge against each other they will stick together for some time, until the sticky compound formed by the jelly on the sperm head is dissolved by the sea water, which occurs after a short time.

This agglutinating effect of the egg-sea water upon the sperm of *Arbacia* gives rise to that ring formation which Lillie considers a proof of positive chemotropism. When a drop of egg-sea water is put into a sufficiently dense suspension of spermatozoa, the spermatozoa at the surface of the drop will agglutinate into practically one dense ring around it, and through the diffusion of some of the dissolved jelly through this ring numerous little clusters will form at the external periphery of the ring, and these clusters will fuse with the ring. In this way the clear region behind the ring originates. The process of fusion continues inside the ring with the result that the latter breaks up into numerous bead-like spherical clusters as Lillie described. In a former paper the writer has pointed out the analogy between the phenomena of transitory sperm agglutination (under the influence of egg-sea water) and surface tension phenomena, inasmuch as two small clusters upon coming in contact fuse into one larger one and inasmuch as elongated clusters break up into two or more spherical clusters.

The ring formation described by Lillie has, therefore,

in the opinion of the writer no connection with positive chemotropism.^d

4. The method of Pfeffer cannot well be used for larger organisms. Barrows²⁵ has devised an apparatus which allowed him to test quantitatively the chemotropic reactions of *Drosophila*. The flies which are positively heliotropic were allowed to go to the light inside of a narrow hollow groove. At a certain spot of the groove two glass bottles were inserted with their openings opposite each other, one of which contained the substance to be tested for chemotropic efficiency, while the other served as a control. The number of flies which on their path were deviated by the bottle containing the substance to be tested were counted and their number compared with that going into the control bottle. The collection of odorous matter in the groove was removed by suction. In this way it was possible to ascertain that the flies are positively chemotropic to ethyl and amyl alcohol, acetic and lactic acid, and to ether. The chemotropic effect of alcohol was increased through the admixture of traces of an ester, e.g., methyl acetate.

In describing the manner of reaction of these flies, Barrows makes the statement that when the odor is weak the fruit fly "attempts first to find the food by the method of trial and error, but as the fly passes into an area of greater stimulation, these movements give way to a direct orientation. This orientation is a well defined tropism response." A similar statement had been made by

^d Lillie also assumes that it is the intensity gradient which determines the direction of motion in tropistic reactions. This is not correct, since positively heliotropic animals go to the light even if by so doing they have to go from strong into weak light (see page 50). The direction of motion in tropistic reactions is determined by differences in the mass of chemical substances on both sides of a symmetrical animal.

Harper for the heliotropism of certain worms, namely that in strong light the animals move by heliotropism, in weak light by "trial and error." These statements are as erroneous as the assertion that while a stone falls under the influence of gravity a feather finds its way down by the method of "trial and error."

Barrows and Harper overlook the rôle of mass action and reaction velocity. When an animal is struck on one side only by light or by a chemically active substance emanating from a center of diffusion, the mass of this substance or of the photochemical reaction product increases on this side. These substances react with some substance of the nerve endings and as soon as the mass of the reaction product reaches a certain quantity the automatic turning, the tropistic reaction, occurs. When the light is strong or when the animal is near the center of diffusion, this happens in a short time and the tropistic character of the reaction is striking, since the animal is quickly put back into its proper orientation if it deviates from it. When the light is weak or when the animal is at some distance from the center of diffusion it will take a longer time before this critical value of the reaction product is reached, and in this case the animal can deviate considerably out of the correct orientation before it is brought back into the right orientation.

CHAPTER XVII

THERMOTROPISM

UNDER the name of thermotropism M. Mendelssohn³⁵²⁻³⁵⁵ has described the observation that *Paramæcia* gather at a definite end of a trough when these ends have a different temperature. The organisms were put into a flat trough resting on tubes through which water was flowing. When the water in the tube had a temperature of 38° at one end of the trough, while the tube at the opposite end was perfused by water of 26° the organisms all gathered at the latter end. If then the temperature of the water in the two tubes was reversed the organisms went to the other end of the trough. If one end had the temperature of 10° the other of 25°, all went to the latter end. In this case we are in all probability not dealing with a tropistic reaction but with a collection of organisms due to the mechanism of motion described for *Paramæcium* by Jennings. When these organisms come suddenly from a region of a moderate temperature to one of lower temperature the activity of their cilia is transitorily reversed, but owing to the asymmetrical arrangement of their cilia they do not go back in the old direction but deviate to one side. This can lead to a collection of *Paramæcia* such as Mendelssohn described.

CHAPTER XVIII

INSTINCTS

THE teleological way of analyzing animal conduct has predominated to such an extent that there has been a tendency to connect all animal reactions with the preservation of the individual and the species. Instincts are considered to be such reactions of the organism as a whole which lead to the nutrition of the individual, the mating of the two sexes, and the care of the offspring. If the tropism theory of animal conduct is justified it must be possible to show that instincts are tropistic reactions.

We have insisted in previous chapters that animals indifferent to light can be made strongly positively or negatively heliotropic by certain chemicals or *vice versa* (e.g., the experiments on certain fresh water crustaceans with acids or alcohol and caffein). We know that the body itself produces at various periods of its existence definite hormones and such hormones can act similarly as the acids or the caffein in the experiments on crustaceans, since it makes no difference whether such substances as acid are introduced into the blood from the outside or from certain tissues of the animal's own body. We know through F. Lillie's observations that in the blood of the male cattle embryo substances circulate which inhibit the development of secondary sexual characters of the female embryo, and we know through Steinach's experiments that the intermediate tissue from the sexual gland of one sex when introduced into the castrated organism of the opposite sex may impart to the latter the sexual instincts of the

former. Hormones produced by definite tissues, therefore, influence the instincts. We want to show that this influence is due to a modification of *tropistic* reactions by the hormones.

Mating in certain fish, like *Fundulus*, consists in the male pressing that part of its body which contains the opening of the sperm duct against the corresponding part of the female body. The latter responds by pressing back, and the pressure of the body is maintained by both sexes through motions of the tail. During this mutual pressure or friction both sexes shed their sexual cells, sperm and eggs, into the water, and since the openings of the cloaca of the male and female, through which the sex cells are shed, are brought almost in contact with each other, sperm and eggs mix at the moment they are shed. This act of mating is due to a stereotropism which exists only during the spawning season and which is supposedly due to certain hormones existing at this time in the animal. The existence of such hormones is also indicated by certain colorations which develop and exist in the male during this period. This stereotropism is to some extent specific since it is exhibited by the contact between the two sexes. The specificity of this stereotropism is of importance and needs further experimental analysis, but that it is in reality a type of common stereotropism is evidenced by the fact that if during the spawning season we keep females isolated from males in an aquarium the females will go through the motions of mating and shed the eggs every time they come in contact with the glass walls of the aquarium. When they are kept permanently isolated from the male they repeat this non-specific purely stereotropic mating throughout the season. The eggs which they shed they quite frequently devour.

These manifestations of a highly developed stereotropism in the segments of the reproductive organs are probably widespread in the animal kingdom. The late Professor Whitman told the writer that male pigeons when kept in isolation will try to go through the motions of mating with any solid object in their field of vision, *e.g.*, glass bottles, and even with objects which give only the optical impression of a solid, namely, their own shadow on the ground.

In ants, the winged males and females become intensely positively heliotropic at the time of mating. Copulation occurs in the air, in the so-called nuptial flight. At a certain time—in the writer's observation toward sunset, when the sky is illuminated at the horizon only—the whole swarm of males and females leave the nest and fly in the direction of the glow. The wedding flight is a heliotropic phenomenon²⁸⁷ presumably due to substances produced in the body during this period. After copulation the female loses its wings and also its positive heliotropism.^a It becomes now intensely stereotropic. When kept in a dark box with pieces of cloth in folds the wingless female will now be found in the folds where its body is as closely as possible in contact with the solids. This positive stereotropism leads the queen to begin a subterranean existence which marks the founding of a new nest. Heliotropism and stereotropism are, therefore, the controlling factors in mating and the starting of a new nest in these ants.²⁸⁷

V. L. Kellogg²⁶⁵ has made observations which show that the nuptial flight in bees is also due to an outburst of positive heliotropism as in the ant.

^a It has already been mentioned that artificial removal of the wings of the fruit fly will also abolish its heliotropism.

In the course of some experiments on the sense-reactions of honey-bees, I have kept a small community of Italian bees in a glass-sided, narrow, high observation hive, so made that any particular bee, marked, which it is desired to observe constantly, can not escape this observation. The hive contains but two frames, one above the other, and is made wholly of glass, except for the wooden frame. It is kept covered, except during observation periods, by a black cloth jacket. The bees live contentedly and normally in this small hive, needing only occasional feeding at times when so many cells are given up for brood that there are not enough left for sufficient stored food supplies. Last spring at the normal swarming time, while standing near the jacketed hive, I heard the excited hum of a beginning swarm and noted the first issuers rushing pell-mell from the entrance. Interested to see the behavior of the community in the hive during such an ecstatic condition as that of swarming, I lifted the cloth jacket, when the excited mass of bees which was pushing frantically down to the small exit in the lower corner of the hive turned with one accord about face and rushed directly upward away from the opening toward and to the top of the hive. Here the bees jammed, struggling violently. I slipped the jacket partly on; the ones covered turned down; the ones below stood undecided; I dropped the jacket completely; the mass began issuing from the exit again; I pulled off the jacket, and again the whole community of excited bees flowed—that is the word for it, so perfectly aligned and so evenly moving were all the individuals of the bee current—up to the closed top of the hive. Leaving the jacket off permanently, I prevented the issuing of the swarm until the ecstasy was passed and the usual quietly busy life of the hive was resumed. About three hours later there was a similar performance and failure to issue from the quickly unjacketed hive. On the next day another attempt to swarm was made, and after nearly an hour of struggling and moving up and down, depending on my manipulation of the black jacket, most of the bees got out of the hive's opening and the swarming came off on a weed bunch near the laboratory. That the issuance from the hive at swarming time depends upon a sudden extra-development of positive heliotropism seems obvious. The ecstasy comes and the bees crowd for the one spot of light in the normal hive, namely, the entrance opening. But when the covering jacket is lifted and the light comes strongly in from above—my hive was under a skylight—they rush toward the top, that is, toward the light. Jacket on and light shut off from above, down they rush; jacket off and light stronger from above than below and they respond like iron filings in front of an electromagnet which has its current suddenly turned on.

Finally there are indications of the rôle of chemotropism in mating. It has been observed for a long time that if a female butterfly is kept hidden from sight in a not too tightly closed box, male butterflies of the same species will be attracted by the box and settle on it. The female apparently gives off a substance to which the male is positively chemotropic. All these observations should be worked out more systematically. The data suffice, however, to indicate that what the biologist and psychologist call instinct are manifestations of tropisms.

The fact that eggs are laid by many insects on material which serves as a nutritive medium for the offspring is a typical instinct. An experimental analysis shows again that the underlying mechanism of the instinct is a positive chemotropism of the mother insect for the type of substance serving her as food; and when the intensity of these volatile substances is very high, *i.e.*, when the insect is on the material, the egg-laying mechanism of the fly is automatically set into motion. Thus the common housefly will deposit its eggs on decaying meat but not on fat; but it will also deposit it on objects smeared over with *asafœtida*, on which the larvæ cannot live. Aseptic banana flies will lay their eggs on sterile banana, although the banana is only an adequate food for the larvæ when yeast grows on it. It seems that the female insect lays her eggs on material for which she is positively chemotropic, and this is generally material which she also eats. The fact that such material serves as food for the coming generation is an accident. Considered in this way, the mystic aspect of the instinctive care of insects for the future generation is replaced by the simple mechanistic conception of a tropistic reaction. In this case natural selection plays a rôle since species whose females would too fre-

quently lay their eggs on material on which the larvæ cannot thrive would be liable to die out.

As an illustration of the rôle of tropisms in the instinctive self-preservation the writer wishes to apologize for selecting an example which he has used so often in previous discussions, namely the rôle of heliotropism in the preservation of the life of the caterpillars of *Porthesia chrysorrhæa*.²⁸⁷ This butterfly lays its eggs upon a shrub, on which the larvæ hatch in the fall and on which they hibernate, as a rule, not far from the ground. As soon as the temperature reaches a certain height, they leave the nest; under natural conditions this happens in the spring when the first leaves have begun to form on the shrub. (The larvæ can, however, be induced to leave the nest at any time in the winter, provided the temperature is raised sufficiently). After leaving the nest, they crawl directly upward on the shrub where they find the leaves on which they feed. If the caterpillars should move down the shrub they would starve, but this they never do, always crawling upward to where they find their food. What gives the caterpillar this never-failing certainty which saves its life and for which the human being might envy the little larva? Is it a dim recollection of experiences of former generations, as Samuel Butler would have us believe? It can be shown that this instinct is merely positive heliotropism and that the light reflected from the sky guides the animals upward. The caterpillars upon waking from their winter sleep are violently positively heliotropic, and it is this heliotropism which makes the animals move upward. At the top of the branch they come in contact with a growing bud and chemical and tactile influences set the mandibles of the young caterpillar into activity. If we put these caterpillars into closed test tubes which lie

with their longitudinal axes at right angles to the window they will all migrate to the window end where they will stay and starve, even if we put their favorite leaves into the test tube close behind them. These larvæ are in this condition slaves of the light.

The few young leaves on top of a twig are quickly eaten by the caterpillar. The light which saved its life by making it creep upward where it finds its food would cause it to starve could the animal not free itself from the bondage of positive heliotropism. After having eaten it is no longer a slave of light but can and does creep downward. It can be shown that a caterpillar after having been fed loses its positive heliotropism almost completely and permanently. If we submit unfed and fed caterpillars of the same nest to the same artificial or natural source of light in two different test tubes the unfed will creep to the light and stay there until they die, while those that have eaten will pay little or no attention to the light. Their positive heliotropism has disappeared and the animal after having eaten can creep in any direction. The restlessness which accompanies the condition of starvation makes the animal leave the top of the branches and creep downward—which is the only direction open to it—where it finds new young leaves on which it can feed. The wonderful hereditary instinct upon which the life of the animal depends is its positive heliotropism in the unfed condition and the loss of this heliotropism after having eaten. The chemical changes following the taking up of the food abolish the heliotropism just as CO_2 arouses positive heliotropism in certain *Daphnia*.

Mayer and Soule have shown that negative geotropism and positive heliotropism keep the caterpillars of *Danais plexippus* on its plant (the milk-weed). The chemical

nature of the leaf starts the eating reactions, but "once the eating reaction be set into play, it tends to continue, so that the larva may then be induced to eat substances which it would never have commenced to eat in the first instance." ³⁵¹

These few examples may suffice to show that the theory of tropisms is at the same time the theory of instincts if due consideration is given to the rôle of hormones in producing certain tropisms and suppressing others. A systematic analysis of instinctive reactions from the viewpoint of the theory of tropisms and hormones will probably yield rich returns. As an example we may quote the fact that diurnal depth migrations of aquatic animals, consisting in an upward motion during the night and a downward motion during the day, are in all probability determined by a periodic change in the sense of heliotropism. ^{183, 300}

CHAPTER XIX

MEMORY IMAGES AND TROPISMS

WHEN a muscle is stimulated several times in succession, the effect of the second or third or later stimulation may be greater than that of the first. A consistently anthropomorphic author should draw the inference that the muscle is gradually learning to react properly. What seems to happen is that the hydrogen ion concentration is raised by the first stimulations to a point where the effect of the stimulation becomes greater. When the stimulations continue and the hydrogen ion concentration becomes still greater, the response of the muscle declines and finally becomes zero; the hydrogen ion concentration has now become too high. The writer observed that when winged plant lice of a *Cineraria* were taken directly from the plant, they did not react as promptly as after they had gone through several heliotropic experiments. There is nothing to indicate that this is a case of "learning," since it may also be the result of a change in the hydrogen ion concentration or of some other reaction product. It may also be the result of some purely mechanical obstacle to rapid locomotion being removed.

We can speak of learning only in such organisms in which the existence of associative memory can be proved. By associative memory we mean that mechanism, by which a stimulus produces not only the direct effects determined by its nature, but also the effects of entirely different stimuli which at some former period by chance attacked the organism at the same time with the given

stimulus. Thus the image or the odor of a rose may call up the memory of persons or surroundings which were present on a former occasion when the image or odor of the flower impressed us. Brain physiology shows that this type of associative memory is the specific function of definite parts of the brain, *e.g.*, the cerebral hemispheres which exist only in definite types of animals. We see also that certain species among vertebrates, insects, crustacea, and cephalopods possess associative memory, while to the knowledge of the writer no adequate proof for its existence has ever been given for worms, starfish, sea urchins, actinians, medusæ, hydroids, or infusorians.²⁹³ Claims for the existence of such memory in these latter groups of animals have frequently been made, but such claims are either plain romance or due to a confusion of reversible physiological processes with the irreversible phenomena of associative memory. The less a scientist is accustomed to rigid quantitative experiments, the more ready he is to confound the reversible after effects of a stimulus—*e.g.*, the after effects due to an increase in hydrogen ion concentration—with indications of associative memory. Learning is only possible where there exists a specific organ of associative memory, the physical mechanism of which is still unknown.

The manifestations of associative memory are generally discussed by the introspective psychologists, who as a rule are not familiar with or do not appreciate the methods of the physicist. There have been made repeated attempts to develop methods for the analysis of associative memory, among which thus far only one satisfies the demands of quantitative science, namely Pawlow's method. As is well known even to the layman, eating causes a flow of saliva. The quantity of saliva excreted

by the parotid (one of the salivary glands) in the dog can be collected and measured. The earlier physiological workers had observed that in a dog which had often been used for the study of the influence of eating upon the flow of saliva, the saliva began to flow whenever the preparations for feeding were made before the eyes of the dog, even when no food was given. Pawlow made use of this fact to study quantitatively the "strength" of such associative phenomena, which he terms "conditioned reflexes" (to escape the terminology and interpretations of the introspective psychologist).⁵³⁷ A fistula^a of the duct of the parotid gland allows the saliva to flow outside the cavity of the mouth. This fistula is connected with a long manometer which by a special air chamber arrangement gives a considerable change in the height of the meniscus for the secretion of as little as one drop of saliva. The variations of the height of the column of liquid in the manometer are observed outside of the room where the dog is. For each dog which is to serve for such experiments the meal is preceded by a certain signal, the sounds of a metronome of definite rhythm, or a definite musical sound, or a definite optical signal, and so forth, which is to form the special conditioned reflex for this dog. After a certain number of repetitions the association is established and from now on the flow of saliva commences from the dog's parotid when the typical signal is given. It was found that the quantity of saliva excreted by the signal changes in a definite sense and quantity when the signal varies or when other conditions accompanying the signal vary.

^a The writer is indebted for the details of Pawlow's method to a short review by Dr. Morgulis.^{537, 538}

Thus in one dog "by persistent training a conditioned reflex has been established to the stimulation with 100 oscillations per minute of the metronome. The stimulation of intermittent sounds of such frequency called forth 6 to 10 drops of saliva every time. The interval between successive oscillations was then modified, the moment of the disappearance of the conditioned salivary reflex indicating the lowest limit of differentiation. Without going into any details of this most interesting investigation or quoting actual data, I will say that the dog could sharply distinguish the shortening of the interval by less than $1/40$ to $1/43$ of a second. Indeed with the well-developed reflex to the stimulation of 100 beats per minute a change of the rate to either 96 or 104 beats was immediately reacted upon by a marked diminution or even complete cessation of the flow of saliva."

This example will give an indication how sensitive is this method of measuring the effect of a memory association.

It is not our purpose to give the details of Pawlow's results—they have only been published in Russian and are therefore not accessible to the writer—but to show that the influence of an associative memory image is as exactly measurable as, *e.g.*, the direct illumination of the eye; and moreover that what we call a memory image is not a "spiritual" but a physical agency. We therefore need not be surprised to find that such memory images or "conditioned reflexes" can vary and multiply the number of possible tropistic reactions.

We have mentioned in the previous chapter that the stereotropism in the mating instinct includes apparently an element of species specificity inasmuch as naturally only males and females of the same species mate. The

late Professor Whitman has shown by experiment that this specificity is, in pigeons at least, not inherited but the effect of memory images (a "conditioned reflex" in the sense of Pawlow). Whitman took the eggs or young of wild species, giving them to the domestic ring-dove to foster, with the result, that the young reared by the ring-doves ever after associated with ring-doves and tried to mate with them. Passenger pigeons when reared by ring-doves refuse to mate with their own species but mate with the species of the foster parents.⁵³⁹ This shows incidentally that racial antagonism is not inherited but acquired.

We have mentioned the fact that the mating instinct is determined by tropisms aroused by specific internal secretions, and that in isolated male pigeons any solid body can arouse the mating reaction. Craig⁵⁴⁰ raised male pigeons in isolation so that they never came in contact with other pigeons until they were adult. One pigeon was hatched in July and isolated in August.

Throughout the autumn and early winter this bird cooed very little. But about the first of February there began a remarkable development of voice and social behavior. The dove was kept in a room where several men were at work, and he directed his display behavior toward these men just as if they belonged to his own species. Each time I put food in his cage he became greatly excited, charging up and down the cage, bowing-and-cooing to me, and pecking my hand whenever it came within his cage. From that day until the day of his death, Jack continued to react in this social manner to human beings. He would bow-and-coo to me at a distance, or to my face when near the cage; but he paid greatest attention to the hand—naturally so, because it was the only part with which he daily came into direct contact. He treated the hand much as if it were a living bird. Not only were his own activities directed toward the hand as if it were a bird, but he received treatment by the hand in the same spirit. The hand could stroke him, preen his neck, even pull the feathers sharply, Jack had absolutely no fear, but ran to the hand to be stroked or teased, showing the joy that all doves show in the attentions of their companions.

When this pigeon was almost a year old it was put into a cage with a female pigeon, but although the female aroused the sexual instinct of the formerly isolated male the latter did not mate with her, but mated with the hand of his attendant when the hand was put into the cage, and this continued throughout the season. Thus the memory images acquired by the bird at an impressionable age and period perverted its sexual tropisms.

It is perhaps of more importance to show that memory images may have a direct orienting influence. The chemotropic phenomenon of an insect laying its egg on a substance which serves as food (for both mother and offspring) and for which the mother is positively chemotropic, may be modified by an act of associative memory, *e.g.*, when a solitary wasp drags the caterpillar on which it lays its eggs to a previously prepared hole in the ground. The essential part of the instinct, the laying of the eggs on the caterpillar, does, perhaps, not differ very much from the fly laying its eggs on decaying meat; and the solitary wasp may be strongly positively chemotropic for the caterpillar on which it lays the eggs, although this has not yet been investigated. But the phenomenon is complicated by a second tropism, which we will call the orienting effect of the memory image. As is well known, the wasp before "going for" the caterpillar digs a hole in the ground to which it afterwards drags the caterpillar, often from a distance. The finding of this previously prepared hole by the returning wasp, the writer would designate as the tropistic or orienting effect of the memory image of the location of this hole; meaning thereby that the memory image of the location of this hole makes the animal return to this location. The conduct of these wasps

is familiar to many readers and the writer may be pardoned for quoting from a formerly published observation.

Ammophila, a solitary wasp, makes a small hole in the ground and then goes out to hunt for a caterpillar, which, when found, it paralyzes by one or several stings. The wasp carries the caterpillar back to the nest, puts it into the hole, and covers the latter with sand. Before this is done, it deposits its eggs on the caterpillar which serves the young larva as food.

An *Ammophila* had made a hole in a flower bed and left the flower bed flying. A little later I saw an *Ammophila* running on the sidewalk of the street in front of the garden, dragging a caterpillar which it held in its mouth. The weight of the caterpillar prevented the wasp from flying. The garden was higher than the sidewalk and separated from it by a stone wall. The wasp repeatedly made an attempt to climb upon the stone wall, but kept falling down. Suspecting that it might have a hole prepared in the garden, I was curious to see whether and how it would find the hole. It followed the wall until it reached the neighboring yard, which had no wall. It now left the street and crawled into this yard, dragging the caterpillar along. Then crawling through the fence which separated the two yards, it dropped the caterpillar near the foot of a tree, and flew away. After a short zigzag flight it alighted on a flower bed in which I noticed two small holes. It soon left the bed and flew back to the tree, not in a straight line but in three stages, stopping twice on its way. At the third stop it landed at the place where the caterpillar lay. The caterpillar was then dragged to the hole, pulled into it, and the hole was covered with tiny stones in the usual way.²⁹³

It is not enough to say that the animal possesses associative memory and returns to the hole; we must add that the brain image of the region of the hole becomes the source of a forced orientation of the animal—of an added special tropism—compelling the animal to return to the region corresponding to the image. And the same may be said in regard to the return of the wasp to the caterpillar which had been temporarily deposited at the foot of the tree.

This example, which might be easily multiplied, will

show the addition necessary to the tropism theory to make it include the endless number of reactions in which associative memory is involved. The psychiatrist would find it easy to supply numerous examples of this type of forced movements toward certain objects which have left a memory image. Since the writer has not investigated this subject sufficiently he is not in a position to give more than a suggestion for the direction of further work. He is inclined to believe that with this enlargement the tropism theory might include human conduct also if we realize that certain memory images may exercise as definite an orienting influence as, *e.g.*, moving retina images or sex hormones.

This tentative extension of the forced movement or tropism theory of animal conduct may explain why higher animals and human beings seem to possess freedom of will, although all movements are of the nature of forced movements. The tropistic effects of memory images and the modification and inhibition of tropisms by memory images make the number of possible reactions so great that prediction becomes almost impossible and it is this impossibility chiefly which gives rise to the doctrine of free will. The theory of free will originated and is held not among physicists but among verbalists. We have shown that an organism goes where its legs carry it and that the direction of the motion is forced upon the organism. When the orienting force is obvious to us, the motion appears as being willed or instinctive; the latter generally when all individuals act alike, machine fashion, the former when different individuals act differently. When a swarm of *Daphnia* is sensitized with CO_2 they all rush to the source of light. This is a machine-like action, and many

will be willing to admit that it is a forced movement or an instinctive reaction. After the CO_2 has evaporated the animals become indifferent to light, and while formerly they had only one degree of freedom of motion they now can move in any direction. In this case the motions appear to be spontaneous or free, since we are not in a position to state why *Daphnia a* moves to the right and *Daphnia b* to the left, etc. As a matter of fact, the motion of each individual is again determined by something but we do not know what it is. The persistent courtship of a human male for a definite individual female may appear as an example of persistent will, yet it is a complicated tropism in which sex hormones and definite memory images are the determining factors. Removal of the sex glands abolishes the courtship and replacing the sex glands of an individual by those of the opposite sex may lead to a complete reversal of the sex instincts. What appears as persistent will action is, therefore, essentially a tropistic reaction. The production of heliotropism by CO_2 in *Daphnia* and the production of the definite courtship of the male *A* for the female *B* are similar phenomena differing only by the nature of the hormones and the additional tropistic effects of certain memory images in the case of courtship. Our conception of the existence of "free will" in human beings rests on the fact that our knowledge is often not sufficiently complete to account for the orienting forces, especially when we carry out a "premeditated" act, or when we carry out an act which gives us pain or may lead to our destruction, and our incomplete knowledge is due to the sheer endless number of possible combinations and mutual inhibitions of the orienting effect of individual memory images.

LITERATURE ^a

- ¹ ABBOTT, J. F., and LIFE, A. C.: Galvanotropism in Bacteria. *Am. J. Physiol.*, 1908, **xxii**, 202-206.
- ² ADAMS, G. P.: On the Negative and Positive Phototropism of the Earthworm *Allolobophora foetida* (Sav.) as Determined by Light of Different Intensities. *Am. J. Physiol.*, 1903, **ix**, 26-34.
- ³ ALLEE, W. C.: An Experimental Analysis of the Relation Between Physiological States and Rheotaxis in Isopoda. *J. Exp. Zool.*, 1912, **xiii**, 269-344.
- ⁴ ALLEE, W. C.: The Effect of Molting on Rheotaxis in Isopods. *Science*, 1913, **xxxvii**, 882-883.
- ⁵ ALLEE, W. C.: Further Studies on Physiological States and Rheotaxis in Isopoda. *J. Exp. Zool.*, 1913, **xv**, 257-295.
- ⁶ ALLEE, W. C.: Certain Relations Between Rheotaxis and Resistance to Potassium Cyanide in Isopoda. *J. Exp. Zool.*, 1914, **xvi**, 397-412.
- ⁷ ALLEE, W. C.: The Ecological Importance of the Rheotactic Reaction of Stream Isopods. *Biol. Bull.*, 1914, **xxvii**, 52-66.
- ⁸ ALLEE, W. C.: Chemical Control of Rheotaxis in *Asellus*. *J. Exp. Zool.*, 1916, **xxi**, 163-198.
- ⁹ ALLEE, W. C., and TASHIRO, S.: Some Relations Between Rheotaxis and the Rate of Carbon Dioxide Production of Isopods. *J. Animal Behav.*, 1914, **iv**, 202-214.
- ¹⁰ ALLEN, G. D.: Reversibility of the Reactions of *Planaria dorotocephala* to a Current of Water. *Biol. Bull.*, 1915, **xxix**, 111-128.
- ¹¹ AREY, L. B.: The Orientation of *Amphioxus* During Locomotion. *J. Exp. Zool.*, 1915, **xix**, 37-44.
- ¹² ARISZ, W. H.: On the Connection Between Stimulus and Effect in Phototropic Curvatures of Seedlings of *Avena sativa*. *Proc. Roy. Acad. Amsterdam*, 1911, **xiii**, 1022-1031.
- ¹³ AXENFELD, D.: Quelques observations sur la vue des arthropodes. *Arch. Ital. Biol.*, 1899, **xxxi**.
- ¹⁴ BACH, H.: Ueber die Abhängigkeit der geotropischen Präsentations- und Reaktionszeit von verschiedenen äusseren Faktoren. *Jahrb. wiss. Bot.*, 1907, **xliv**, 57-123.
- ¹⁵ BALSS, H.: Über die Chemorezeption bei Garneelen. *Biol. Centr.*, 1913, **xxxiii**, 508-512.

^a The list of literature does not claim to be complete. Aside from unintentional omissions, some of the controversial and amateurish publications have not been included.

- 16 BANCROFT, F. W.: Note on the Galvanotropic Reactions of the *Mедуsa Polyorchis penicillata*, A. Agassiz. *J. Exp. Zool.*, 1904, i, 289-292.
- 17 BANCROFT, F. W.: Ueber die Gültigkeit des Pflüger'schen Gesetzes für die galvanotropischen Reaktionen von *Paramæcium*. *Arch. ges. Physiol.*, 1905, cvii, 535-556.
- 18 BANCROFT, F. W.: On the Influence of the Relative Concentration of Calcium Ions on the Reversal of the Polar Effects of the Galvanic Current in *Paramæcium*. *J. Physiol.*, 1906, xxxiv, 444-463.
- 19 BANCROFT, F. W.: The Control of Galvanotropism in *Paramæcium* by Chemical Substances. *Univ. Cal. Pub. Physiol.*, 1906, iii, 21-31.
- 20 BANCROFT, F. W.: The Mechanism of Galvanotropic Orientation in *Volvox*. *J. Exp. Zool.*, 1907, iv, 157-163.
- 21 BANCROFT, F. W.: Heliotropism, Differential Sensibility, and Galvanotropism in *Euglena*. *J. Exp. Zool.*, 1913, xv, 383-428.
- 22 BANTA, A. M.: Experiments on the Light and Tactile Reactions of a Cave Variety and an Open Water Variety of an Amphipod Species. *Proc. Soc. Exp. Biol. and Med.*, 1913, x, 192.
- 23 BARANETZKI, J.: Influence de la lumière sur les plasmodia des myxomycètes. *Mém. Soc. Sc. Nat. Cherbourg*, 1876, xix, 321-360.
- 24 BARRATT, J. O. W.: Der Einfluss der Konzentration auf die Chemotaxis. *Z. allg. Physiol.*, 1905, v, 73-94.
- 25 BARROWS, W. M.: The Reactions of the Pomace Fly, *Drosophila ampelophila* Loew, to Odorous Substances. *J. Exp. Zool.*, 1907, iv, 515-537.
- 26 BAUER, V.: Ueber die reflektorische Regulierung der Schwimmbewegungen bei den Mysiden, mit besonderer Berücksichtigung der doppelsinnigen Reizbarkeit der Augen. *Z. allg. Physiol.*, 1908, viii, 343-370.
- 27 BAUER, V.: Ueber sukzessiven Helligkeitskontrast bei Fischen. *Centr. Physiol.*, 1909, xxiii, 593-599.
- 28 BAUER, V.: Ueber das Farbenunterscheidungsvermögen der Fische. *Arch. ges. Physiol.*, 1910, cxxxiii, 7-26.
- 29 BAUER, V.: Zur Kenntnis der Lebensweise von *Pecten jacobæus* L. Im besonderen über die Funktion der Augen. *Zool. Jahrb. Abt. allg. Zool.*, 1912, xxxiii, 127-150.
- 30 BAUNACKE, W.: Statische Sinnesorgane bei den Nepiden. *Zool. Jahrb. Abt. Anat.*, 1912-13, xxxiv, 179-346.
- 31 BAUNACKE, W.: Studien zur Frage nach der Statocystenfunktion. I. Statische Reflexe bei Mollusken. *Biol. Centr.*, 1913, xxxiii, 427-452.

- ³² BAUNACKE, W.: II. Noch einmal die Geotaxis unserer Mollusken. *Biol. Centr.*, 1914, xxxiv, 371-385; 497-523.
- ³³ BEER, TH.: Vergleichend-physiologische Studien zur Statocysten-funktion. I. Ueber den angeblichen Gehörsinn und das angebliche Gehörorgan der Crustaceen. *Arch. ges. Physiol.*, 1898, lxxiii, 1-41.
- ³⁴ BEER, TH.: II. Versuche an Crustaceen (*Penaeus membranaceus*). *Arch. ges. Physiol.*, 1899, lxxiv, 361-382.
- ³⁵ BENGT, J.: Der richtende Einfluss strömenden Wassers auf wachsende Pflanzen und Pflanzenteile (Rheotropismus). *Ber. bot. Ges.*, 1883, i, 512-521.
- ³⁶ BERNSTEIN, J.: Chemotropische Bewegung eines Quecksilbertropfens. Zur Theorie der amöboiden Bewegung. *Arch. ges. Physiol.*, 1900, lxxx, 628-637.
- ³⁷ BERT, P.: Les animaux voient-ils les mêmes rayons lumineux que nous? *Mém. Soc. Sc. Phys. et Nat. Bordeaux*, 1868, vi, 375-383.
- ³⁸ BERT, P.: Sur la question de savoir si tous les animaux voient les mêmes rayons lumineux que nous. *Arch. de Physiol.*, 1869, ii, 547-554.
- ³⁹ BETHE, A.: Ueber die Erhaltung des Gleichgewichts. *Biol. Centr.*, 1894, xiv, 95-114; 563-582.
- ⁴⁰ BETHE, A.: Die Otocyste von Mysis. *Zool. Jahrb. Abt. Anat.*, 1895, viii, 544-564.
- ⁴¹ BETHE, A.: Die Locomotion des Haifisches (*Scyllium*) und ihre Beziehungen zu den einzelnen Gehirnteilen und zum Labyrinth. *Arch. ges. Physiol.*, 1899, lxxvi, 470-493.
- ⁴² BIRGE, E. A.: The Vertical Distribution of the Limnetic Crustacea of Lake Mendota. *Biol. Centr.*, 1897, xvii, 371-374.
- ⁴³ BIRUKOFF, B.: Untersuchungen über Galvanotaxis. *Arch. ges. Physiol.*, 1899, lxxvii, 555-585.
- ⁴⁴ BIRUKOFF, B.: Zur Theorie der Galvanotaxis. *Arch. Anat. u. Physiol., Physiol. Abt.*, 1904, 271-296.
- ⁴⁵ BIRUKOFF, B.: Zur Theorie der Galvanotaxis. II. *Arch. ges. Physiol.*, 1906, cxi, 95-143.
- ⁴⁶ BLAAUW, A. H.: The Intensity of Light and the Length of Illumination in the Phototropic Curvature in Seedlings of *Avena sativa* (Oats). *Proc. Roy. Akad. Amsterdam*, 1908.
- ⁴⁷ BLAAUW, A. H.: Die Perzeption des Lichtes. *Rec. trav. bot. Néerlandais*, 1909, v, 209-377.
- ⁴⁸ BLAAUW, A. H.: Licht und Wachstum. *Z. Bot.*, 1914, vi, 641-703; 1915, vii, 465-532.
- ^{48a} BLASIUS, E., and SCHWEIZER, F.: Elektrotropismus und verwandte Erscheinungen. *Arch. ges. Physiol.*, 1893, liii, 493-543.

- 49 BOHN, G.: Les *Convoluta roscoffensis* et la théorie des causes actuelles. *Bull. Mus. Paris*, 1903, 352-364.
- 50 BOHN, G.: Théorie nouvelle du phototropisme. *Compt. rend. Acad. Sc.*, 1904, cxxxix, 890-891.
- 51 BOHN, G.: Attractions et oscillations des animaux marins sous l'influence de la lumière. Recherches nouvelles relatives au phototactisme et au phototropisme. *Mém. Inst. génér. Psychol.*, 1905, i, 1-111.
- 52 BOHN, G.: Impulsions motrices d'origine oculaire chez les crustacés. (Deuxième mémoire relatif au phototactisme et au phototropisme.) *Bull. Inst. génér. Psychol.*, 1905, v, 412-454.
- 53 BOHN, G.: Intervention des réactions oscillatoires dans les tropismes. *Compt. rend. Assoc. Française avancement des Sc., Congrès de Reims*, 1907, 700-706.
- 54 BOHN, G.: Observations biologiques sur le branchellion de la torpille. *Bull. Station biol. Arcachon*, 1907, x, 283-296.
- 55 BOHN, G.: Les tropismes, la sensibilité différentielle et les associations chez le branchellion de la torpille. *Compt. rend. Soc. Biol.*, 1907, lxiii, 545-548.
- 56 BOHN, G.: A propos des lois de l'excitabilité par la lumière. I. Le retour progressif à l'état d'immobilité, après une stimulation mécanique. *Compt. rend. Soc. Biol.*, 1907, lxiii, 655-658.
- 57 BOHN, G.: II. Du changement de signe du phototropisme en tant que manifestation de la sensibilité différentielle. *Compt. rend. Soc. Biol.*, 1907, lxiii, 756-759.
- 58 BOHN, G.: Introduction à la psychologie des animaux à symétrie rayonnée. I. Les états physiologiques des actinies. *Bull. Inst. génér. Psychol.*, 1907, vii, 81-129; 135-182.
- 59 BOHN, G.: II. Les essais et erreurs chez les étoiles de mer et les ophiures. *Bull. Inst. génér. Psychol.*, 1908, viii, 21-102.
- 60 BOHN, G.: Les rythmes vitaux chez les actinies. *Compt. rend. Assoc. Française avancement des Sc.*, 1908, 613.
- 61 BOHN, G.: De l'orientation chez les patelles. *Compt. rend. Acad. Sc.*, 1909, cxlviii, 868-870.
- 62 BOHN, G.: Les variations de la sensibilité périphérique chez les animaux. *Bull. Sc. France et Belgique*, 1909, xliii, 481-519.
- 63 BOHN, G.: Quelques problèmes généraux relatifs à l'activité des animaux inférieurs. *Bull. Inst. génér. Psychol.*, 1909, ix, 439-466.
- 64 BOHN, G.: Quelques observations sur les chenilles des dunes. *Bull. Inst. génér. Psychol.*, 1909, ix, 543-549.
- 65 BOHN, G.: La naissance de l'intelligence. Paris, 1909.

- ⁶⁶ BOHN, G.: Les tropismes. *Rapport VI^{me} Congr. Internat. Psychol. Genève*, 1909, pp. 15.
- ⁶⁷ BOHN, G.: A propos les lois de l'excitabilité par la lumière. III. De l'influence de l'éclairement du fond sur le signe des réactions vis-à-vis la lumière. *Compt. rend. Soc. Biol.*, 1909, lxvi, 18-20.
- ⁶⁸ BOHN, G.: IV. Sur les changements périodiques du signe des réactions. *Compt. rend. Soc. Biol.*, 1909, lxvii, 4-6.
- ⁶⁹ BOHN, G.: V. Intervention de la vitesse des réactions chimiques dans la désensibilisation par la lumière. *Compt. rend. Soc. Biol.*, 1910, lxviii, 1114-1117.
- ⁷⁰ BOHN, G.: La sensibilisation et la désensibilisation des animaux. *Compt. rend. Assoc. Française avancement des Sc., Congrès de Toulouse*, 1910, 214-222.
- ⁷¹ BOHN, G.: Quelques expériences de modification des réactions chez les animaux, suivies de considérations sur les mécanismes chimiques de l'évolution. *Bull. Sc. France et Belgique*, 1911, xlv, 217-238.
- ⁷² BOHN, G.: La nouvelle psychologie animale. Paris, 1911, pp. 200.
- ⁷³ BOHN, G.: La sensibilité des animaux aux variations de pression. *Compt. rend. Acad. Sc.*, 1912, cliv, 240-242.
- ⁷⁴ BOHN, G.: Les variations de la sensibilité en relation avec les variations de l'état chimique interne. *Compt. rend. Acad. Sc.*, 1912, cliv, 388-391.
- ⁷⁵ BOHN, G.: L'étude des phénomènes mnémiques chez les organismes inférieurs. *J. Psychol. u. Neurol.*, 1913, xx, 199-209.
- ⁷⁶ BORING, E. G.: Note on the Negative Reaction Under Light Adaptation in the *Planarian*. *J. Animal Behav.*, 1912, ii, 229-248.
- ⁷⁷ BORN, G.: Biologische Untersuchungen. Ueber den Einfluss der Schwere auf das Froschei. *Arch. mikr. Anat.*, 1885, xxiv, 475.
- ⁷⁸ BREUER, J.: Ueber die Funktion der Bogengänge des Ohrlabyrinths. *Med. Jahrb.*, 1874.
- ⁷⁹ BREUER, J.: Beiträge zur Lehre vom statischen Sinne. *Med. Jahrb.*, 1875.
- ⁸⁰ BREUER, J.: Ueber die Funktion der Otolithenapparate. *Arch. ges. Physiol.*, 1891, xlviii, 195-306.
- ^{80a} BREUER, J.: Ueber den Galvanotropismus (Galvanotaxis) bei Fischen. *Sitzungsb. Akad. Wiss. Wien. mathem.-naturw. Kl.*, 1905, cxiv, 27-56.
- ^{80b} BREUER, J., and KREIDL, A.: Ueber die scheinbare Drehung des Gesichtsfeldes, während der Einwirkung einer Centrifugalkraft. *Arch. ges. Physiol.*, 1898, lxx, 491-510.
- ⁸¹ BRUCHMANN, H.: Chemotaxis der *Lycopodium*-Spermatozoiden. *Flora*, 1908-09, xcix, 193-202.

- ⁸² BRUN, R.: Die Raumorientierung der Ameisen und das Orientierungsproblem im allgemeinen. Jena, 1914, pp. 242.
- ⁸³ BRUNDIN, T. M.: Light Reactions of Terrestrial Amphipods. *J. Animal Behav.*, 1913, iii, 334-352.
- ⁸⁴ v. BUDDENBROCK, W.: Untersuchungen über die Schwimmbewegungen und die Statocysten der Gattung *Pecten*. *Sitzgsb. Heidelberger Akad. Wiss., mathem.-naturw. Kl.*, 1911, pp. 24.
- ⁸⁵ v. BUDDENBROCK, W.: Ueber die Funktion der Statocysten im Sande grabender Meerestiere (*Arenicola* und *Synapta*). *Biol. Centr.*, 1912, xxxii, 564-585.
- ⁸⁶ v. BUDDENBROCK, W.: Ueber die Funktion der Statocysten von *Branchiomma vesiculosum*. *Verhandl. naturhist.-med. Vereines, Heidelberg*, 1913, N.F. xii, 256-261.
- ⁸⁷ v. BUDDENBROCK, W.: Ueber die Orientierung der Krebse im Raum. *Zool. Jahrb. Abt. Zool.*, 1914, xxxiv, 479-514.
- ⁸⁸ v. BUDDENBROCK, W.: A Criticism of the Tropism Theory of Jacques Loeb. *J. Animal Behav.*, 1916, vi, 341-366.
- ⁸⁹ BULLER, A. H. R.: Contributions to Our Knowledge of the Physiology of the Spermatozoa of Ferns. *Annals Bot.*, 1900, xiv, 543-582.
- ⁹⁰ BULLER, A. H. R.: Is Chemotaxis a Factor in the Fertilization of the Eggs of Animals? *Quart. J. Micr. Sc.*, 1902-03, xlv, 145-176.
- ⁹¹ BOYSEN-JENSEN, P.: Ueber die Leitung des phototropischen Reizes in der *Avenakoleoptile*. *Ber. bot. Ges.*, 1913, xxxi, 559-566.
- ⁹² BUNTING, M.: Ueber die Bedeutung der Otolithenorgane für die geotropischen Funktionen von *Astacus fluviatilis*. *Arch. ges. Physiol.*, 1893, liv, 531-537.
- ⁹³ CARLGREN, O.: Der Galvanotropismus und die innere Kataphorese. *Z. allg. Physiol.*, 1905, v, 123-130.
- ⁹⁴ CARLGREN, O.: Ueber die Einwirkung des konstanten galvanischen Stromes auf niedere Organismen. *Arch. Anat. u. Physiol., Physiol. Abt.*, 1900, 49-76.
- ⁹⁵ CARPENTER, F. W.: The Reactions of the Pomace Fly (*Drosophila ampelophila*, Loew) to Light, Gravity, and Mechanical Stimulation. *Am. Nat.*, 1905, xxxix, 157-171.
- ^{95a} CLAPARÈDE, E.: Les tropismes devant la psychologie. *J. Psychol. u. Neurol.*, 1908, xiii, 150-160.
- ⁹⁶ CLARK, G. P.: On the Relation of the Otocysts to Equilibrium Phenomena in *Gelasimus pugilator* and *Platyonichus ocellatus*. *J. Physiol.*, 1896, xix, 327-343.
- ⁹⁷ CLARK, O. L.: Ueber negativen Phototropismus bei *Avena sativa*. *Z. Bot.*, 1913, v, 737-770.

- ⁹⁸ COEHN, A., and BARRATT, W.: Ueber Galvanotaxis vom Standpunkte der physikalischen Chemie. *Z. allg. Physiol.*, 1905, v, 1-9.
- ⁹⁹ COHN, F.: Ueber die Gesetze der Bewegung mikroskopischer Tiere und Pflanzen unter Einfluss des Lichtes. *Jahr.-ber. Schles. Ges. vaterl. Kultur*, 1864, xlii, 35-36.
- ¹⁰⁰ COLE, L. J.: The Influence of Direction vs. Intensity of Light in Determining the Phototropic Responses of Organisms. *Science*, 1907, xxv, 784.
- ¹⁰¹ CONGDON, E. D.: Recent Studies Upon the Locomotor Responses of Animals to White Light. *J. Comp. Neurol. and Psychol.*, 1908, xviii, 309-328.
- ¹⁰² CORNETZ, V.: Ueber den Gebrauch des Ausdruckes "tropisch" und über den Charakter der Richtungskraft bei Ameisen. *Arch. ges. Physiol.*, 1912, cxlvii, 215-233.
- ¹⁰³ COWLES, R. P.: Stimuli Produced by Light and by Contact with Solid Walls as Factors in the Behavior of Ophiuroids. *J. Exp. Zool.*, 1910, ix, 387-416.
- ¹⁰⁴ COWLES, R. P.: Reaction to Light and Other Points in the Behavior of the Starfish. *Papers from Tortugas Lab. Carnegie Inst. Washington*, 1911, iii, 95-110.
- ¹⁰⁵ COWLES, R. P.: The Influence of White and Black Walls on the Direction of Locomotion of the Starfish. *J. Animal Behav.*, 1914, iv, 380-382.
- ^{105a} CRAIG, W.: The Voices of Pigeons Regarded as a Means of Social Control. *Am. J. Sociology*, 1908, xiv, 86-100.
- ^{105b} CRAIG, W.: Male Doves Reared in Isolation. *J. Animal Behav.*, 1914, iv, 121-133.
- ^{105c} CRAIG, W.: Appetites and Aversions as Constituents of Instincts. *Biol. Bull.*, 1918, xxxiv, 97-107.
- ¹⁰⁶ CROZIER, W. J.: The Orientation of a Holothurian by Light. *Am. J. Physiol.*, 1914, xxxvi, 8-20.
- ¹⁰⁷ CROZIER, W. J.: The Behavior of Holothurians in Balanced Illumination. *Am. J. Physiol.*, 1917, xliii, 510-513.
- ¹⁰⁸ CROZIER, W. J.: The Photoreceptors of *Amphioxus*. *Anat. Rec.*, 1917, xi, 520.
- ^{108a} CROZIER, W. J.: The Photic Sensitivity of *Balanoglossus*. *J. Exp. Zool.*, 1917, xxiv, 211-217.
- ¹⁰⁹ CZAPEK, F.: Ueber Zusammenwirken von Heliotropismus und Geotropismus. *Sitzungsber. Akad. Wiss. Wien. mathem.-naturw. Kl.*, 1895, civ.
- ¹¹⁰ CZAPEK, F.: Untersuchungen über Geotropismus. *Jahrb. wiss. Bot.*, 1895, xxvii, 243-339.

- 111 CZAPPEK, F.: Weitere Beiträge zur Kenntnis der geotropischen Reizbewegungen. *Jahrb. wiss. Bot.*, 1898, xxxii, 175-308.
- 112 DALE, H. H.: Galvanotaxis and Chemotaxis of Ciliate Infusoria. *J. Physiol.*, 1901, xxvi, 291-361.
- 113 DAVENPORT, C. B.: Experimental Morphology. Part I. Effects of Chemical and Physical Agents Upon Protoplasm. New York, 1897.
- 114 DAVENPORT, C. B., and CANNON, W. B.: On the Determination of the Direction and Rate of Movement of Organisms by Light. *J. Physiol.*, 1897, xxi, 22-32.
- 115 DAVENPORT, C. B., and LEWIS, F. T.: Phototaxis of *Daphnia*. *Science*, 1899, ix, 368.
- 116 DAVENPORT, C. B., and PERKINS, H.: A Contribution to the Study of Geotaxis in the Higher Animals. *J. Physiol.*, 1897, xxii, 99-110.
- 117 DAY, E. C.: The Effect of Colored Light on Pigment Migration in the Eye of the Crayfish. *Bull. Mus. Comp. Zool.*, 1911, liii, 303-343.
- 118 DELAGE, Y.: Étude expérimentale sur les illusions statiques et dynamiques de direction pour servir à déterminer les fonctions des canaux semicirculaires de l'oreille interne. *Arch. Zool. expér. et génér.*, 1886, (2) iv.
- 119 DELAGE, Y.: Sur une fonction nouvelle des otocystes comme organes d'orientation locomotrice. *Arch. Zool. expér. et génér.*, 1887, (2) v, 1-26.
- 120 DEWITZ, J.: Ueber die Vereinigung der Spermatozoen mit dem Ei. *Arch. ges. Physiol.*, 1885, xxxvii, 219-223.
- 121 DEWITZ, J.: Ueber Gesetzmässigkeit in der Ortsveränderung der Spermatozoen und in der Vereinigung derselben mit dem Ei. *Arch. ges. Physiol.*, 1886, xxxviii, 358-385.
- 122 DEWITZ, J.: Ueber den Rheotropismus bei Tieren. *Arch. Physiol.*, 1899 (Suppl.), 231-244.
- 123 DOLLEY, W. L., JR.: Reactions to Light in *Vanessa antiopa*, with Special Reference to Circus Movements. *J. Exp. Zool.*, 1916, xx, 357-420.
- 124 DRIESCH, H.: Heliotropismus bei Hydroidpolypen. *Zool. Jahrb.*, 1890, v, 147-156.
- 125 DRIESCH, H.: Die taktische Reizbarkeit der Mesenchymzellen von *Echinus microtuberculatus*. *Arch. Entwicklungsmech.*, 1896, iii, 362-380.
- 126 DRIESCH, H.: Die organischen Regulationen. Leipzig, 1901, pp. 228.
- 127 DUBOIS, R.: Sur le mécanisme des fonctions photodermatique et photogénique dans le siphon du *Pholas dactylus*. *Compt. rend. Acad. Sc.*, 1889, cix, 233-235.

- ¹²⁸ DUBOIS, R.: Sur l'action des agents modificateurs de la contraction photodermatique chez le *Pholas dactylus*. *Compt. rend. Acad. Sc.*, 1898, cix, 320-322.
- ¹²⁹ DUBOIS, R.: Sur la perception des radiations lumineuses par la peau, chez les Protées aveugles des grottes de la Carniole. *Compt. rend. Acad. Sc.*, 1890, cx, 358-361.
- ¹³⁰ DUBOIS, R.: Note sur l'action de la lumière sur les échinodermes (oursin). *Commun. 9me. Cong. internat. Zool.*, Monaco, 1913, (1), 8-9.
- ¹³¹ DUSTIN, A. P.: Le rôle des tropismes et de l'odogenèse dans la régénération du système nerveux. *Arch. Biol.*, 1910, xxv, 269-388.
- ¹³² ENGELMANN, T. W.: Ueber Reizung kontraktile Protoplasmas durch plötzliche Beleuchtung. *Arch. ges. Physiol.*, 1879, xix, 1-7.
- ¹³³ ENGELMANN, T. W.: Ueber Licht- und Farbenperzeption niederster Organismen. *Arch. ges. Physiol.*, 1882, xxix, 387-400.
- ¹³⁴ ENGELMANN, T. W.: *Bacterium photometricum*. Ein Beitrag zur vergleichenden Physiologie des Licht- und Farbensinnes. *Arch. ges. Physiol.*, 1882, xxx, 95-124.
- ¹³⁵ ENGELMANN, T. W.: Ueber die Funktion der Otolithen. *Zool. Anz.*, 1887, x, 591, 664.
- ¹³⁶ ENGELMANN, T. W.: Die Purpurbakterien und ihre Beziehungen zum Licht. *Bot. Ztg.*, 1888, xvi, 661-669, 677-689, 693-701, 709-720.
- ¹³⁷ ENGLISCH, E.: Ueber die Wirkung intermittierender Belichtungen auf Bromsilbergelatine. *Arch. wiss. Phot.*, 1899, i, 117-131.
- ¹³⁸ ENGLISCH, E.: Ueber den zeitlichen Verlauf der durch das Licht verursachten Veränderungen der Bromsilbergelatine. *Arch. wiss. Phot.*, 1900, ii, 131-134.
- ¹³⁹ ERHARD, H.: Beitrag zur Kenntnis des Lichtsinnes der Daphniden. *Biol. Centr.*, 1913, xxxiii, 494-496.
- ¹⁴⁰ ESTERLY, C. O.: The Reactions of *Cyclops* to Light and Gravity. *Am. J. Physiol.*, 1907, xviii, 47-57.
- ¹⁴¹ EWALD, J. R.: Physiologische Untersuchungen über das Endorgan des Nervus octavus. Wiesbaden, 1892.
- ¹⁴² EWALD, J. R.: Ueber die Wirkung des galvanischen Stroms bei der Längsdurchströmung ganzer Wirbeltiere. *Arch. ges. Physiol.*, 1894, iv, 606-621 (Berichtigung, 1894, lvi, 354).
- ¹⁴³ EWALD, W. E.: Ueber Orientierung, Lokomotion und Lichtreaktionen einiger Cladoceren und deren Bedeutung für die Theorie der Tropismen. *Biol. Centr.*, 1910, xxx, 1-16, 49-63, 379-399.
- ¹⁴⁴ EWALD, W. E.: On Artificial Modification of Light Reactions and the Influence of Electrolytes on Phototaxis. *J. Exp. Zool.*, 1912, xiii, 591-612.

- 145 EWALD, W. E.: The Applicability of the Photochemical Energy Law to Light Reactions in Animals. *Science*, 1913, xxxviii, 236-237.
- 146 EWALD, W. E.: Ist die Lehre vom tierischen Phototropismus widerlegt? *Arch. Entwicklungsmech.*, 1913, xxxvii, 581-598.
- 147 EWALD, W. E.: Versuche zur Analyse der Licht- und Farbenreaktionen eines Wirbellosen (*Daphnia pulex*). *Z. Sinnesphysiol.*, 1914, xlviii, 285-324.
- 148 EYCLESHYMER, A. C.: The Reactions to Light of the Decapitated Young *Necturus*. *J. Comp. Neurol. and Psychol.*, 1908, xviii, 303-308.
- 149 FAUVEL, P., and BOHN, G.: Le rythme des marées chez les diatomées littorales. *Compt. rend. Soc. Biol.*, 1907, lxii, 121-123.
- 150 FIGDOR, W.: Ueber Helio- und Geotropismus der Gramineenblätter. *Ber. bot. Ges.*, 1905, xxiii, 182-191.
- 151 FIGDOR, W.: Experimentelle Studien über die heliotropische Empfindlichkeit der Pflanzen. *Wiesner Festschrift, Wien*, 1908.
- 152 FIGDOR, W.: Heliotropische Reizleitung bei Begonia-Blättern. *Ann. Jardin bot. Buitenzorg.*, 1910 (Suppl.), iii, 453-460.
- 153 FIGDOR, W.: Ueber thigmotropische Empfindlichkeit der Asparagus-Sprosse. *Sitznsgb. Akad. Wiss. Wien. mathem.-naturw. Kl. Abt. I*, 1915, cxxiv, 353.
- 154 FITTING, H.: Untersuchungen über den geotropischen Reizvorgang. *Jahrb. wiss. Bot.*, 1905, xli, 221-398.
- 155 FLOURENS, P.: Recherches expérimentales sur les propriétés et les fonctions du système nerveux dans les animaux vertébrés. Paris, 1842, pp. xxviii + 516.
- 156 FORSSMAN, J.: Ueber die Ursachen, welche die Wachstumsrichtung der peripheren Nervenfasern bei der Regeneration bestimmen. *Beitr. path. Anat.*, 1898, xxiv, 56-100.
- 157 FORSSMAN, J.: Zur Kenntnis des Neurotropismus. *Beitr. path. Anat.*, 1900, xxvii, 407-430.
- 158 FRANDSEN, P.: Studies on the Reactions of *Limax maximus* to Directive Stimuli. *Proc. Am. Acad. Arts and Sc.*, 1901, xxxvii, 185-227.
- 159 FRANZ, V.: Phototaxis und Wanderung. Nach Versuchen mit Jungfischen und Fischlarven. *Int. Rev. ges. Hydrobiol. u. Hydrographie*, 1910, iii, 306-334.
- 160 FRANZ, V.: Beiträge zur Kenntnis der Phototaxis. Nach Versuchen an Süßwassertieren. *Int. Rev. ges. Hydrobiol. u. Hydrographie, Biol. Suppl. (2)*, 1911, 1-11.
- 161 FRANZ, V.: Weitere Phototaxisstudien. I. Zur Phototaxis bei Fischen. II. Phototaxis bei marinen Crustaceen. III. Phototaktische Lokomotionsperioden bei *Hemimysis*. *Int. Rev. ges. Hydrobiol. u. Hydrographie, Biol. Suppl. (3)*, 1911, 1-23.

- 162 FRANZ, V.: Zur Frage der vertikalen Wanderungen der Planktontiere. *Arch. Hydrobiol. u. Planktonkunde*, 1912, vii, 493-499.
- 163 FRANZ, V.: Die phototaktischen Erscheinungen im Tierreiche und ihre Rolle im Freileben der Tiere. *Zool. Jahrb.*, 1913, xxxiii, 259-286.
- 164 v. FRISCH, K.: Ueber farbige Anpassung bei Fischen. *Zool. Jahrb.*, 1912, xxxii, 171-230.
- 165 v. FRISCH, K.: Sind die Fische farbenblind? *Zool. Jahrb.*, 1912, xxxiii, 107-126.
- 166 v. FRISCH, K.: Ueber die Farbenanpassung des *Crenilabrus*. *Zool. Jahrb.*, 1912, xxxiii, 151-164.
- 167 v. FRISCH, K.: Weitere Untersuchungen über den Farbensinn der Fische. *Zool. Jahrb.*, 1913, xxxiv, 43-68.
- 168 v. FRISCH, K.: Der Farbensinn und Formensinn der Biene. *Zool. Jahrb.*, 1914, xxxv, 1-182.
- 169 v. FRISCH, K., and KUPELWIESER, H.: Ueber den Einfluss der Lichtfarbe auf die phototaktischen Reaktionen niederer Krebse. *Biol. Centr.*, 1913, xxxiii, 517-552.
- 170 FRÖHLICH, F. W.: Vergleichende Untersuchungen über den Licht- und Farbensinn. *Deutsch. med. Wchnschr.*, 1913, xxxix, 1453-1456.
- 171 FRÖSCHEL, P.: Untersuchung über die heliotropische Präsentationszeit. I. *Sitzgsb. Akad. Wiss. Wien. mathem.-naturw. Kl.*, 1908, cxvii, 235-256.
- 172 FRÖSCHEL, P.: Untersuchung über die heliotropische Präsentationszeit. II. *Sitzgsb. Akad. Wiss. Wien. mathem.-naturw. Kl.*, 1909, cxviii, 1247-1294.
- 173 FUCHS, R. F.: Der Farbenwechsel und die chromatische Hautfunktion der Tiere. *Winterstein's Handb. vergl. Physiol.*, 1914, iii, I. Hälfte 2, 1189-1656.
- 174 GALLIANO, E. F.: Beitrag zur Untersuchung der Chemotaxis der *Paramecien*. *Z. allg. Physiol.*, 1914, xvi, 359-372.
- 175 GARREY, W. E.: The Effect of Ions Upon the Aggregation of Flagellated Infusoria. *Am. J. Physiol.*, 1900, iii, 291-315.
- 176 GARREY, W. E.: A Sight Reflex Shown by Sticklebacks. *Biol. Bull.*, 1905, viii, 79-84.
- 177 GARREY, W. E.: Proof of the Muscle Tension Theory of Heliotropism. *Proc. Nat. Acad. Sc.*, 1917, iii, 602-609.
- 178 GOLTZ, F.: Ueber die Verrichtungen des Grosshirns. I-V. *Arch. ges. Physiol.*, 1876, xiii, 1-44; 1877, xiv, 412-443; 1879, xx, 1-54; 1881, xxvi, 1-49; 1884, xxxiv, 450-505.
- 179 GRABER, V.: Fundamentalversuche über die Helligkeits- und Farbeempfindlichkeit augenloser und geblendeter Tiere. *Sitzgsb. Akad. Wiss. Wien*, 1883, lxxxvii, 201-236.

- 180 GRABER, V.: Grundlinien zur Erforschung des Helligkeits- und Farbensinnes der Tiere. Leipzig, 1884, pp. vii + 322.
- 181 GRABER, V.: Ueber die Helligkeits- und Farbenempfindlichkeit einiger Meertiere. *Sitzungsber. Akad. Wiss. Wien.*, 1885, xci.
- 182 GRABER, V.: Thermische Experimente an der Küchenschabe (*Periplaneta orientalis*). *Arch. ges. Physiol.*, 1887, xli, 240-256.
- 183 GROOM, T. T., and LOEB, J.: Der Heliotropismus der Nauplien von *Balanus perforatus* und die periodischen Tiefenwanderungen pelagischer Tiere. *Biol. Centr.*, 1890, x, 160-177.
- 184 GROSS, A. O.: The Reactions of Arthropods to Monochromatic Lights of Equal Intensities. *J. Exp. Zool.*, 1913, xiv, 467-514.
- 185 HABERLANDT, G.: Ueber die Perzeption des geotropischen Reizes. *Ber. bot. Ges.*, 1900, xviii, 261-272.
- 186 HADLEY, P. B.: The Relation of Optical Stimuli to Rheotaxis in the American Lobster (*Homarus americanus*). *Am. J. Physiol.*, 1906, xvii, 326-343.
- 187 HADLEY, P. B.: Galvanotaxis in Larvæ of the American Lobster (*Homarus americanus*). *Am. J. Physiol.*, 1907, xix, 39-52.
- 188 HADLEY, P. B.: The Reaction of Blinded Lobsters to Light. *Am. J. Physiol.*, 1908, xxi, 180-199.
- 189 HADLEY, P. B.: Reactions of Young Lobsters Determined by Food Stimuli. *Science*, 1912, xxxv, 1000-1002.
- 190 HARPER, E. H.: Reactions to Light and Mechanical Stimuli in the Earthworm, *Perichæta bermudensis* (Beddard). *Biol. Bull.*, 1905, x, 17-34.
- 191 HARPER, E. H.: Tropic and Shock Reactions in *Perichæta* and *Lumbricus*. *J. Comp. Neurol. and Psychol.*, 1909, xix, 569-587.
- 192 HARPER, E. H.: The Geotropism of *Paramæcium*. *J. Morphol.*, 1911, xxii, 993-1000.
- 193 HARPER, E. H.: Magnetic Control of Geotropism in *Paramæcium*. *J. Animal Behav.*, 1912, ii, 181-189.
- 194 HARRINGTON, N. R., and LEAMING, E.: The Reaction of *Amœba* to Lights of Different Colors. *Am. J. Physiol.*, 1899, iii, 9-18.
- 195 HASEMAN, J. D.: The Rhythmical Movements of *Littorina littorea* Synchronous with Ocean Tides. *Biol. Bull.*, 1911, xxi, 113-121.
- 196 HAUSMANN, W.: Die photodynamische Wirkung des Chlorophylls und ihre Beziehung zur photosynthetischen Assimilation der Pflanzen. *Jahrb. wiss. Bot.*, 1909, xlv, 599-623.
- 197 HELMHOLTZ, H.: Handbuch der physiologischen Optik. Hamburg, 1909-11, 3. Ed.
- 198 HENRI, MME. V., and HENRI, V.: Excitation des organismes par les rayons ultra-violet. *Compt. rend. Soc. Biol.*, 1912, lxxii, 992-996; lxxiii, 326-327.

- 199 HENRI, V., and LARGUIER DES BANCELS, J.: Photochimie de la rétine. *J. Physiol. et Path. génér.*, 1911, xiii, 841-856.
- 200 HENRI, V., and LARGUIER DES BANCELS, J.: Un nouveau type de temps de réaction. *Compt. rend. Soc. Biol.*, 1912, lxxiii, 55-56.
- 201 HENRI, V., and LARGUIER DES BANCELS, J.: L'excitation provoquée par les rayons ultra-violetes comparée avec les excitations visuelle et nerveuse, d'une part, et les réactions photochimiques, d'autre. Lois des phénomènes. *Compt. rend. Soc. Biol.*, 1912, lxxiii, 328-329.
- 202 HENRI, V., and LARGUIER DES BANCELS, J.: Sur l'interprétation des lois de Weber et de Jost: recherches sur les réactions des *Cyclops* exposées à la lumière ultra-violette. *Arch. Psychol.*, 1912, xii, 329-342.
- 203 HERBST, C.: Ueber die Bedeutung der Reizphysiologie für die kausale Auffassung von Vorgängen in der tierischen Ontogenese. I. *Biol. Centr.*, 1894, xiv, 657-666, 689-697, 727-744, 753-771, 800-810.
- 204 HERMANN, L.: Eine Wirkung galvanischer Ströme auf Organismen. *Arch. ges. Physiol.*, 1885, xxxvii, 457-460.
- 205 HERMANN, L.: Weitere Untersuchungen über das Verhalten der Froschlarven im galvanischen Strom. *Arch. ges. Physiol.*, 1886, xxxix, 414-419.
- 205a HERMANN, L., and MATTHIAS, F.: Der Galvanotropismus der Larven von *Rana temporaria* und der Fische. *Arch. ges. Physiol.*, 1894, lvii, 391-405.
- 206 HERMS, W. B.: The Photic Reactions of Sacrophagid Flies, Especially *Lucilia caesar* Linn. and *Calliphora vomitoria* Linn. *J. Exp. Zool.*, 1911, x, 167-226.
- 207 HERTEL, E.: Ueber die Beeinflussung des Organismus durch Licht, speciell durch die chemisch wirksamen Strahlen. *Z. allg. Physiol.*, 1904, iv, 1-43.
- 208 HERTEL, E.: Ueber physiologische Wirkung von Strahlen verschiedener Wellenlänge. *Z. allg. Physiol.*, 1905, v, 95-122.
- 209 HESS, C.: Untersuchungen über den Lichtsinn bei Fischen. *Arch. Augenheilk.*, 1909, lxiv, 1-38.
- 210 HESS, C.: Untersuchungen über den Lichtsinn bei wirbellosen Tieren. *Arch. Augenheilk.*, 1909, lxiv, 39-61.
- 211 HESS, C.: Neue Untersuchungen über den Lichtsinn bei wirbellosen Tieren. *Arch. ges. Physiol.*, 1910, cxxxvi, 282-367.
- 212 HESS, C.: Experimentelle Untersuchungen zur vergleichenden Physiologie des Gesichtssinnes. *Arch. ges. Physiol.*, 1911, cxlii, 405-446.
- 213 HESS, C.: Der Gesichtssinn. *Winterstein's Handb. vergl. Physiol.*, 1912, iv, 555-840.
- 214 HESS, C.: Neue Untersuchungen zur vergleichenden Physiologie des Gesichtssinnes. *Zool. Jahrb. Abt. Zool.*, 1913, xxxiii, 387-440.

- 215 HESS, C.: Experimentelle Untersuchungen über den angeblichen Farbensinn der Bienen. *Zool. Jahrb. Abt. Zool.*, 1913, xxxiv, 81-106.
- 216 HESS, C.: Eine neue Methode zur Untersuchung des Lichtsinnes bei Krebsen. *Arch. vergl. Ophthalmol.*, 1913-14, iv, 52-67.
- 217 HESS, C.: Untersuchungen über den Lichtsinn mariner Würmer und Krebse. *Arch. ges. Physiol.*, 1914, clv, 421-435.
- 218 HESS, C.: Untersuchungen über den Lichtsinn bei Echinodermen. *Arch. ges. Physiol.*, 1914, clx, 1-26.
- 219 HESS, C.: Messende Untersuchung des Lichtsinnes der Biene. *Arch. ges. Physiol.*, 1916, clxiii, 289-320.
- 220 HESSE, R.: Untersuchungen über die Organe der Lichtempfindung bei niederen Tieren. I. Die Organe der Lichtempfindung bei den Lubriciden. *Z. wiss. Zool.*, 1896, lxi, 393-419.
- 221 HESSE, R.: II. Die Augen der Plathelminthen, insonderheit der trikladen Turbellarien. *Z. wiss. Zool.*, 1897, lxii, 527-582.
- 222 HESSE, R.: IV. Die Sehorgane des *Amphioxus*. *Z. wiss. Zool.*, 1898, lxiii, 456-464.
- 223 HESSE, R.: Die Lichtempfindung des *Amphioxus*. *Anat. Anz.*, 1898, xiv, 556.
- 224 HÖGYES, A.: Der Nervenmechanismus der assoziierten Augenbewegungen. I-II. *Mitt. mathem.-naturw. Kl. Ungar. Akad. Wiss. Budapest*, 1881, x, 1-62; xi, 1-100. (Ref. *Biol. Centr.*, 1881-82, i, 216-220.)
- 225 HOLMES, S. J.: Phototaxis in the Amphipoda. *Am. J. Physiol.*, 1901, v, 211-234.
- 226 HOLMES, S. J.: Phototaxis in *Volvox*. *Biol. Bull.*, 1903, iv, 319-326.
- 227 HOLMES, S. J.: The Selection of Random Movements as a Factor in Phototaxis. *J. Comp. Neurol. and Psychol.*, 1905, xv, 98-112.
- 228 HOLMES, S. J.: The Reactions of *Ranatra* to Light. *J. Comp. Neurol. and Psychol.*, 1905, xv, 305-349.
- 229 HOLMES, S. J.: Observations on the Young of *Ranatra quadridentata* Stal. *Biol. Bull.*, 1907, xii, 158-164.
- 230 HOLMES, S. J.: Phototaxis in Fiddler Crabs and Its Relation to Theories of Orientation. *J. Comp. Neurol. and Psychol.*, 1908, xviii, 493-497.
- 231 HOLMES, S. J.: Description of a New Species of *Eubbranchipus* from Wisconsin with Observations on Its Reaction to Light. *Trans. Wis. Acad. Sc., Arts and Letters*, 1910, xvi, pt. II, 1252-1255.
- 232 HOLMES, S. J.: Pleasure, Pain and the Beginnings of Intelligence. *J. Comp. Neurol. and Psychol.*, 1910, xx, 145-164.
- 233 HOLMES, S. J.: Evolution of Animal Intelligence. New York, 1911, pp. 296.

- 234 HOLMES, S. J.: The Reactions of Mosquitoes to Light in Different Periods of Their History. *J. Animal Behav.*, 1911, i, 29-32.
- 235 HOLMES, S. J.: The Beginnings of Intelligence. *Science*, 1911, xxxiii, 473-480.
- 236 HOLMES, S. J.: The Tropisms and Their Relation to More Complex Modes of Behavior. *Bull. Wis. Nat. Hist. Soc.*, 1912, x, 13-23.
- 237 HOLMES, S. J.: Phototaxis in the Sea Urchin, *Arbacia punctulata*. *J. Animal Behav.*, 1912, ii, 126-136.
- 238 HOLMES, S. J.: Studies in Animal Behavior. Boston, 1916, pp. 266.
- 239 HOLMES, S. J., and MCGRAW, K. W.: Some Experiments on the Method of Orientation to Light. *J. Animal Behav.*, 1913, iii, 367-373.
- 240 HOLT, E. B., and LEE, F. S.: The Theory of Phototactic Response. *Am. J. Physiol.*, 1901, iv, 460-481.
- 241 HOWARD, L. O.: Butterflies Attracted to Light at Night. *Proc. Ent. Soc., Washington*, 1889, iv.
- 242 ILYIN, P.: Das Gehörbläschen als Gleichgewichtsorgan bei den Pterotracheidae. *Centr. Physiol.*, 1900, xiii, 691-694.
- 243 JACKSON, H. H. T.: The Control of Phototactic Reactions in *Hyalella* by Chemicals. *J. Comp. Neurol. and Psychol.*, 1910, xx, 259-263.
- 244 JENNINGS, H. S.: Studies on Reactions to Stimuli in Unicellular Organisms. I. Reactions to Chemical, Osmotic and Mechanical Stimuli in the Ciliate Infusoria. *J. Physiol.*, 1897, xxi, 258-322.
- 245 JENNINGS, H. S.: II. The Mechanism of the Motor Reactions of *Paramecium*. *Am. J. Physiol.*, 1899, ii, 311-341.
- 246 JENNINGS, H. S.: III. Reactions to Localized Stimuli in *Spirostomum* and *Stentor*. *Am. Nat.*, 1899, xxxiii, 373-389.
- 247 JENNINGS, H. S.: V. On the Movements and Motor Reflexes of the Flagellata and Ciliata. *Am. J. Physiol.*, 1900, iii, 229-260.
- 248 JENNINGS, H. S.: VI. On the Reactions of *Chilomonas* to Organic Acids. *Am. J. Physiol.*, 1900, iii, 397-403.
- 249 JENNINGS, H. S., and CROSBY, J. H.: VII. The Manner in Which Bacteria React to Stimuli, Especially to Chemical Stimuli. *Am. J. Physiol.*, 1901, vi, 31-37.
- 250 JENNINGS, H. S., and MOORE, E. H.: VIII. On the Reactions of Infusoria to Carbonic and Other Acids, with Especial Reference to the Causes of the Gatherings Spontaneously Formed. *Am. J. Physiol.*, 1902, vi, 233-250.
- 251 JENNINGS, H. S.: Contributions to the Study of the Behavior of Lower Organisms. *Carnegie Institution of Washington*, Pub. No. 16, 1904, pp. 256, 81 figs.

- 252 JENNINGS, H. S.: Modifiability in Behavior. II. Factors Determining Direction and Character of Movement in the Earthworm. *J. Exp. Zool.*, 1906, iii, 435-455.
- 253 JENNINGS, H. S.: Behavior of the Lower Organisms. New York, 1906, pp. xiv + 366.
- 254 JENNINGS, H. S.: The Interpretation of the Behavior of the Lower Organisms. *Science*, 1908, xxvii, 698-710.
- 255 JENNINGS, H. S.: Tropisms. *Rapport VI^{me} Congr. Internat. Psychol. Genève*, 1909, pp. 20.
- 255a JENSEN, P.: Ueber den Geotropismus niederer Organismen. *Arch. ges. Physiol.*, 1893, liii, 428-480.
- 256 JORDAN, H.: Rheotropic Responses of *Epinephelus striatus* Bloch. *Am. J. Physiol.*, 1917, xliii, 438-454.
- 257 JORDAN, H.: Integumentary Photosensitivity in a Marine Fish, *Epinephelus striatus* Bloch. *Am. J. Physiol.*, 1917, xlii, 259-274.
- 258 JUDAY, C.: The Diurnal Movement of Plancton Crustacea. *Trans. Wis. Acad. Sc., Arts and Letters*, 1904, xiv, 534-568.
- 259 KAFKA, G.: Einführung in die Tierpsychologie. I. Die Sinne der Wirbellosen. Leipzig, 1914, xii+594.
- 260 KANDA, S.: On the Geotropism of *Paramæcium* and *Spirostomum*. *Biol. Bull.*, 1914, xxvi, 1-24.
- 261 KANDA, S.: The Reversibility of the Geotropism of *Arenicola* Larvæ by Salts. *Am. J. Physiol.*, 1914, xxxv, 162-176.
- 262 KANDA, S.: Geotropism in Animals. *Am. J. Psychol.*, 1915, xxvi, 417-427.
- 263 KANDA, S.: Studies on the Geotropism of the Marine Snail, *Littorina littorea*. *Biol. Bull.*, 1916, xxx, 57-84.
- 264 KANDA, S.: The Geotropism of Freshwater Snails. *Biol. Bull.*, 1916, xxx, 85-97.
- 264a KANDA, S.: Further Studies on the Geotropism of *Paramæcium caudatum*. *Biol. Bull.*, 1918, xxxiv, 108-119.
- 265 KELLOGG, V. L.: Some Insect Reflexes. *Science*, 1903, xviii, 693-696.
- 266 KELLOGG, V. L.: Some Silkworm Moth Reflexes. *Biol. Bull.*, 1907, xii, 152-154.
- 267 KNIEP, H.: Untersuchungen über die Chemotaxis von Bakterien. *Jahrb. wiss. Bot.*, 1906, xliii, 215-270.
- 268 KRANICHFELD, H.: Zum Farbensinn der Bienen. *Biol. Centr.*, 1915, xxxv, 39-46.
- 269 KRECKER, F. H.: Phenomena of Orientation Exhibited by Ephemeridæ. *Biol. Bull.*, 1915, xxix, 381-388.
- 270 KREIDL, A.: Weitere Beiträge zur Physiologie des Orlabyrinthes. *Sitzungsab. Akad. Wiss. Wien, mathem.-naturw. Kl.*, 1892, ci, 469-480; 1893, cii, 149-174.

- 271 KRIBS, H. G.: The Reactions of *Æolosoma* to Chemical Stimuli. *J. Exp. Zool.*, 1910, viii, 43-74.
- 272 KÜHNE, W.: Untersuchungen über das Protoplasma und die Kontractilität. Leipzig, 1864, pp. 158.
- 273 KÜHNE, W.: Chemische Vorgänge in der Netzhaut. *Hermann's Handb. Physiol.*, 1879, iii, pt. 1, 235-342.
- 274 KUSANO, S.: Studies on the Chemotactic and Other Related Reactions of the Swarmspores of Myxomycetes. *J. Coll. Agriculture, Imp. Univ. Tokyo*, 1909, ii.
- 275 LASAREFF, P.: Ionentheorie der Nerven- und Muskelreizung. *Arch. ges. Physiol.*, 1910, cxxxv, 196-204.
- 276 LAURENS, H.: The Reactions of Amphibians to Monochromatic Lights of Equal Intensity. *Bull. Mus. Comp. Zool.*, 1911, xliii, 253-302.
- 277 LAURENS, H.: The Reactions of Normal and Eyeless Amphibian Larvæ to Light. *J. Exp. Zool.*, 1914, xvi, 195-210.
- 278 LEE, F. S.: A Study of the Sense of Equilibrium in Fishes. I. *J. Physiol.*, 1893, xv, 311-348.
- 279 LEE, F. S.: A Study of the Sense of Equilibrium in Fishes. II. *J. Physiol.*, 1894-95, xvii, 192-210.
- 280 LIDFORSS, B.: Ueber den Chemotropismus der Pollenschläuche. *Ber. bot. Ges.*, 1899, xvii, 236-242.
- 281 LIDFORSS, B.: Ueber die Reizbewegungen der *Marchantia*-Spermatozoiden. *Jahrb. wiss. Bot.*, 1905, xli, 65-87.
- 282 LIDFORSS, B.: Ueber die Chemotaxis eines Thiospirillum. *Ber. bot. Ges.*, 1912, xxx, 262-274.
- 283 LILLIE, F. R.: Studies of Fertilization. V. The Behavior of the Spermatozoa of *Nereis* and *Arbacia* with Special Reference to Egg-extractives. *J. Exp. Zool.*, 1913, xiv, 515-574.
- 284 LOEB, J.: Beiträge zur Physiologie des Grosshirns. *Arch. ges. Physiol.*, 1886, xxxix, 265-346.
- 285 LOEB, J.: Die Orientierung der Tiere gegen das Licht (tierischer Heliotropismus). *Sitzgsb. Würzb. physik.-med. Ges.*, 1888.
- 286 LOEB, J.: Die Orientierung der Tiere gegen die Schwerkraft der Erde (tierischer Geotropismus). *Sitzgsb. Würzb. physik.-med. Ges.*, 1888.
- 287 LOEB, J.: Der Heliotropismus der Tiere und seine Uebereinstimmung mit dem Heliotropismus der Pflanzen. Würzburg, 1889, pp. 118.
- 288 LOEB, J.: Weitere Untersuchungen über den Heliotropismus der Tiere und seine Uebereinstimmung mit dem Heliotropismus der Pflanzen. (Heliotropische Krümmungen bei Tieren). *Arch. ges. Physiol.*, 1890, xlvii, 391-416.

- 289 LOEB, J.: Ueber Geotropismus bei Tieren. *Arch. ges. Physiol.*, 1891, xlix, 175-189.
- 290 LOEB, J.: Ueber den Anteil des Hörnerven an den nach Gehirnverletzung auftretenden Zwangsbewegungen, Zwangslagen und assoziierten Stellungsänderungen der Bulbi und Extremitäten. *Arch. ges. Physiol.*, 1891, l, 66-83.
- 290a LOEB, J.: Untersuchungen zur physiologischen Morphologie der Tiere. I. Heteromorphose. II. Organbildung und Wachstum. Würzburg, 1891-1892.
- 291 LOEB, J.: Ueber künstliche Umwandlung positiv heliotropischer Tiere in negativ heliotropische und umgekehrt. *Arch. ges. Physiol.*, 1893, liv, 81-107.
- 292 LOEB, J.: Zur Theorie der physiologischen Licht- und Schwerkraftwirkungen. *Arch. ges. Physiol.*, 1897, lxvi, 439-466.
- 293 LOEB, J.: Comparative Physiology of the Brain and Comparative Psychology. New York, 1900, x+309.
- 294 LOEB, J.: Studies in General Physiology. Chicago, 1905, 2 vols., x+782.
- 295 LOEB, J.: The Dynamics of Living Matter. New York, 1906, xi+233.
- 296 LOEB, J.: Ueber die Erregung von positivem Heliotropismus durch Säure, insbesondere Kohlensäure und von negativem Heliotropismus durch ultraviolette Strahlen. *Arch. ges. Physiol.*, 1906, cxv, 564-581.
- 297 LOEB, J.: Concerning the Theory of Tropisms. *J. Exp. Zool.*, 1907, iv, 151-156.
- 298 LOEB, J.: Ueber die Summation heliotropischer und geotropischer Wirkungen bei den auf der Drehscheibe ausgelösten kompensatorischen Kopfbewegungen. *Arch. ges. Physiol.*, 1907, cxvi, 368-374.
- 299 LOEB, J.: Chemische Konstitution und physiologische Wirksamkeit von Alkoholen und Säuren. II. *Biochem. Z.*, 1909, xxiii, 93-96.
- 300 LOEB, J.: Die Tropismen. *Winterstein's Handb. vergl. Physiol.*, 1911, iv, 451-519.
- 301 LOEB, J.: The Mechanistic Conception of Life. Chicago, 1912, pp. 232.
- 302 LOEB, J.: On the Nature of the Conditions Which Determine or Prevent the Entrance of the Spermatozoon into the Egg. *Am. Nat.*, 1915, xlix, 257-285.
- 303 LOEB, J.: The Organism as a Whole. From a Physico-chemical Viewpoint. New York, 1916, pp. 379.

- 303a LOEB, J.: The Chemical Basis of Regeneration and Geotropism. *Science*, 1917, xlv, 115-118.
- 303b LOEB, J.: Influence of the Leaf upon Root Formation and Geotropic Curvature in the Stem of *Bryophyllum calycinum* and the Possibility of a Hormone Theory of These Processes. *Bot. Gaz.*, 1917, lxiii, 25-50.
- 303c LOEB, J.: The Chemical Mechanism of Regeneration. *Ann. Inst. Pasteur*, 1918, xxxii, 1-16.
- 304 LOEB, J., and BUDGETT, S. P.: Zur Theorie des Galvanotropismus. IV. Ueber die Ausscheidung electropositiver Ionen an der äusseren Anodenfläche protoplasmatischer Gebilde als Ursache der Abweichungen vom Pflüger'schen Erregungsgesetz. *Arch. ges. Physiol.*, 1897, lxxv, 518-534.
- 305 LOEB, J., and EWALD, W. F.: Ueber die Gültigkeit des Bunsen-Roscoe'schen Gesetzes für die heliotropische Erscheinung bei Tieren. *Centr. Physiol.*, 1914, xxvii, 1165-1168.
- 306 LOEB, J., and GARREY, W. E.: Zur Theorie des Galvanotropismus. II. Versuche an Wirbeltieren. *Arch. ges. Physiol.*, 1896, lxxv, 41-47.
- 307 LOEB, J., and MAXWELL, S. S.: Zur Theorie des Galvanotropismus. *Arch. ges. Physiol.*, 1896, lxiii, 121-144.
- 308 LOEB, J., and MAXWELL, S. S.: Further Proof of the Identity of Heliotropism in Animals and Plants. *Univ. Cal. Pub. Physiol.*, 1910, iii, 195-197.
- 309 LOEB, J., and NORTHPROP, J. H.: Heliotropic Animals as Photometers, on the Basis of the Validity of the Bunsen-Roscoe Law for Heliotropic Reactions. *Proc. Nat. Acad. Sc.*, 1917, iii, 539-544.
- 310 LOEB, J., and WASTENEYS, H.: On the Identity of Heliotropism in Animals and Plants. *Proc. Nat. Acad. Sc.*, 1915, i, 44-47; *Science*, 1915, xli, 328-330.
- 311 LOEB, J., and WASTENEYS, H.: The Relative Efficiency of Various Parts of the Spectrum for the Heliotropic Reactions of Animals and Plants. *J. Exp. Zool.*, 1915, xix, 23-35; 1916, xx, 217-236.
- 312 LOEB, J., and WASTENEYS, H.: A Re-examination of the Applicability of the Bunsen-Roscoe Law to the Phenomena of Animal Heliotropism. *J. Exp. Zool.*, 1917, xxii, 187-192.
- 313 LÖHNER, L.: Untersuchungen über den sogenannten Totstellreflex der Arthropoden. *Z. allg. Physiol.*, 1914, xvi, 373-418.
- 314 LUBBOCK, J.: On the Sense of Color Among Some of the Lower Animals. I and II. *J. Linn. Soc. (Zool.)*, 1881, xvi, 121-127; 1882, xvii, 205-214.
- 315 LUBBOCK, J.: On the Senses, Instincts and Intelligence of Animals, with Special Reference to Insects. Internat. sc. Series, London, 1899.

- ³¹⁶ LUBBOCK, J.: Ants, Bees and Wasps. New York, 1904, xiii+435.
- ³¹⁷ LUDLOFF, K.: Untersuchungen über den Galvanotropismus. *Arch. ges. Physiol.*, 1895, lix, 525-554.
- ³¹⁸ LYON, E. P.: The Functions of the Otocyst. *J. Comp. Neurol. and Psychol.*, 1898, viii, 238-245.
- ³¹⁹ LYON, E. P.: A Contribution to the Comparative Physiology of Compensatory Motions. *Am. J. Physiol.*, 1899, iii, 86-114.
- ³²⁰ LYON, E. P.: Compensatory Motions in Fishes. *Am. J. Physiol.*, 1900, iv, 77-82.
- ³²¹ LYON, E. P.: On Rheotropism. I. Rheotropism in Fishes. *Am. J. Physiol.*, 1904, xii, 149-161.
- ³²² LYON, E. P.: Rheotropism in Fishes. *Biol. Bull.*, 1905, viii, 238-239.
- ³²³ LYON, E. P.: On the Theory of Geotropism in *Paramæcium*. *Am. J. Physiol.*, 1905, xiv, 421-432.
- ³²⁴ LYON, E. P.: Note on the Geotropism of *Arbacia* Larvæ. *Biol. Bull.*, 1906, xii, 21-22.
- ³²⁵ LYON, E. P.: Note on the Heliotropism of *Palæmonetes* Larvæ. *Biol. Bull.*, 1906, xii, 23-25.
- ³²⁶ LYON, E. P.: On Rheotropism. II. Rheotropism of Fish Blind in One Eye. *Am. J. Physiol.*, 1909, xxiv, 244-251.
- ^{326a} LYON, E. P.: Note on the Geotropism of *Paramæcium*. *Biol. Bull.*, 1918, xxxiv, 120.
- ^{326b} MCCLENDON, J. F.: Protozoan Studies. *J. Exp. Zool.*, 1909, vi, 265-283.
- ³²⁷ MACCURDY, H.: Some Effects of Sunlight in the Starfish. *Science*, 1913, xxxvi, 98-100.
- ^{327a} MCEWEN, R. S.: The Reactions to Light and to Gravity in *Drosophila* and its Mutants. *J. Exp. Zool.*, 1918, xxv, 49-106.
- ³²⁸ MCGINNIS, M. O.: Reactions of *Branchipus serratus* to Light, Heat and Gravity. *J. Exp. Zool.*, 1911, x, 227-240.
- ³²⁹ MACH, E.: Physikalische Versuche über den Gleichgewichtssinn des Menschen. *Sitzungsab. Akad. Wiss. Wien.*, 1873, lxviii; 1874, lxix.
- ³³⁰ MACH, E.: Grundlinien der Lehre von den Bewegungsempfindungen. Leipzig, 1875, pp. 127.
- ³³¹ MACH, E.: Beiträge zur Analyse der Empfindungen. Jena, 1902.
- ³³² MAGNUS, R.: Welche Teile des Zentralnervensystems müssen für das Zustandekommen der tonischen Hals- und Labyrinthreflexe auf die Körpermuskulatur vorhanden sein? *Arch. ges. Physiol.*, 1914, clxx, 224-250.
- ³³³ MAGNUS, R., and DE KLEIJN, A.: Die Abhängigkeit des Tonus der Extremitätenmuskeln von der Kopfstellung. *Arch. ges. Physiol.*, 1912, cxlv, 455-548.

- ³³⁴ MAGNUS, R., and DE KLEIJN, A.: Die Abhängigkeit des Tonus der Nackenmuskeln von der Kopfstellung. *Arch. ges. Physiol.*, 1912, cxlvii, 403-416.
- ³³⁵ MAGNUS, R., and DE KLEIJN, A.: Die Abhängigkeit der Körperstellung vom Kopfstande beim normalen Kaninchen. *Arch. ges. Physiol.*, 1913, cliv, 163-177.
- ³³⁶ MAGNUS, R., and DE KLEIJN, A.: Analyse der Folgezustände einseitiger Labyrinthexstirpation mit besonderer Berücksichtigung der Rolle der tonischen Halsreflexe. *Arch. ges. Physiol.*, 1913, cliv, 178-306.
- ³³⁷ MAGNUS, R., and VAN LEEUWEN, W. S.: Die akuten und die dauernden Folgen des Ausfalles der tonischen Hals- und Labyrinthreflexe. *Arch. ges. Physiol.*, 1914, cliv, 157-217.
- ³³⁸ MAGNUS, R., and WOLF, C. G. L.: Weitere Mitteilungen über den Einfluss der Kopfstellung auf den Gliedertonus. *Arch. ges. Physiol.*, 1913, cxlix, 447-461.
- ³³⁹ MARCHAL, P.: Le retour au nid chez le *Pompilus sericeus* V. d. L. *Compt. rend. Soc. Biol.*, 1900, lii, 1113-1115.
- ³⁴⁰ MASSART, J.: Recherches sur les organismes inférieurs. I. La loi du Weber vérifiée pour l'héliotropisme du champignon. *Bull. Acad. Roy. Belg.*, 1888, (3) xvi, 590.
- ³⁴¹ MASSART, J.: Sur l'irritabilité des spermatozoides dans l'oeuf de la grenouille. *Bull. Acad. Roy. Belg.*, 1888, (3) xv; 1889, xviii.
- ³⁴² MASSART, J.: La sensibilité tactile chez les organismes inférieurs. *J. Soc. Roy. Sc. med. et nat., Bruxelles*, 1890.
- ³⁴³ MASSART, J.: Recherches sur les organismes inférieurs. III. La sensibilité à la gravitation. *Bull. Acad. Roy. Belg.*, 1891, (3) xxii, 158-167.
- ³⁴⁴ MASSART, J.: Essai de classification des réflexes non-nerveux. *Ann. Inst. Pasteur*, 1901, xv, 635-672.
- ³⁴⁵ MASSART, J.: Versuch einer Einteilung der nichtnervösen Reflexe. *Biol. Centr.*, 1902, xxii, 9-23.
- ³⁴⁶ MAST, S. O.: Light and the Behavior of Organisms. New York. 1911, pp. 410+xi.
- ³⁴⁷ MAST, S. O.: Behavior of Fire-flies (*Photinus pyralis*?) with Special Reference to the Problem of Orientation. *J. Animal Behav.*, 1912, ii, 256-272.
- ³⁴⁸ MAST, S. O.: The Relation between Spectral Color and Stimulation in the Lower Organisms. *J. Exp. Zool.*, 1917, xxii, 471-528.
- ^{348a} MATULA, J.: Untersuchungen über die Funktionen des Zentralnervensystems bei Insekten. *Arch. ges. Physiol.*, 1911, cxxxviii, 388-456.

- 349 MAXWELL, S. S.: Beiträge zur Gehirnphysiologie der Anneliden. *Arch. ges. Physiol.*, 1897, lxvii, 263-297.
- 350 MAXWELL, S. S.: Experiments on the Functions of the Internal Ear. *Univ. Cal. Pub. Physiol.*, 1910, iv, 1-4.
- 351 MAYER, A. G., and SOULE, C. G.: Some Reactions of Caterpillars and Moths. *J. Exp. Zool.*, 1906, iii, 415-433.
- 352 MENDELSSOHN, M.: Ueber den Thermotropismus einzelliger Organismen. *Arch. ges. Physiol.*, 1895, ix, 1-27.
- 353 MENDELSSOHN, M.: Recherches sur la thermotaxie des organismes unicellulaires. *J. Physiol. et Path. génér.*, 1902, iv, 393-409.
- 354 MENDELSSOHN, M.: Recherches sur l'interférence de la thermotaxie avec d'autres tactismes et sur le mécanisme du mouvement thermotactique. *J. Physiol. et Path. génér.*, 1902, iv, 475-488.
- 355 MENDELSSOHN, M.: Quelques considérations sur la nature et le rôle biologique de la thermotaxie. *J. Physiol. et Path. génér.*, 1902, iv, 489-496.
- 356 MENKE, H.: Periodische Bewegungen und ihr Zusammenhang mit Licht und Stoffwechsel. *Arch. ges. Physiol.*, 1911, cxl, 37-91.
- 357 MEREJKOWSKY, C. DE: Les crustacés inférieurs distinguent-ils les couleurs? *Compt. rend. Acad. Sc.*, 1881, xciii, 1160-1161.
- 358 MILLER, F. R.: Galvanotropism in the Crayfish. *J. Physiol.*, 1907, xxxv, 215-229.
- 359 MINKIEWICZ, R.: Sur le chromotropisme et son inversion artificielle. *Compt. rend. Acad. Sc.*, 1906, cxliii, 785-787.
- 360 MINKIEWICZ, R.: Le rôle des phénomènes chromotropiques dans l'étude des problèmes biologiques et psycho-physiologiques. *Compt. rend. Acad. Sc.*, 1906, cxliii, 934-935.
- 361 MINKIEWICZ, R.: Une expérience sur la nature du chromotropisme chez les némeres. *Compt. rend. Acad. Sc.*, 1912, clv, 229-231.
- 362 MITSUKURI, K.: Negative Phototaxis and Other Properties of *Littorina* as Factors in Determining Its Habitat. *Annotationes Zoologicae Japonenses*, 1901, iv, 1-19.
- 363 MOLISCH, H.: Untersuchungen über den Hydrotropismus. *Sitzungsber. Akad. Wiss. Wien. mathem.-naturw. Kl.*, 1883.
- 364 MOORE, ANNE: Some Facts Concerning Geotropic Gatherings of *Paramacia*. *Am. J. Physiol.*, 1903, ix, 238-244.
- 365 MOORE, A. R.: On the Righting Movements of the Starfish. *Biol. Bull.*, 1910, xix, 235-239.
- 366 MOORE, A. R.: Concerning Negative Phototropism in *Daphnia pulex*. *J. Exp. Zool.*, 1912, xiii, 573-575.
- 367 MOORE, A. R.: Negative Phototropism in *Diaptomus* by Means of Strychnine. *Univ. Cal. Pub. Physiol.*, 1912, iv, 185-186.

- 368 MOORE, A. R.: The Negative Phototropism of *Diaptomus* Through the Agency of Caffein, Strychnine, and Atropin. *Science*, 1913, xxxviii, 131-133.
- 369 MOORE, A. R.: The Mechanism of Orientation in *Gonium*. *J. Exp. Zool.*, 1916, xxi, 431-432.
- 369a MOORE, A. R.: The Action of Strychnine on Certain Invertebrates. *J. Pharm. and Exp. Therap.*, 1916, ix, 167-169.
- 370 MOORE, A. R., and KELLOGG, F. M.: Note on the Galvanotropic Response of the Earthworm. *Biol. Bull.*, 1916, xxx, 131-134.
- 371 MOORE, B.: Observations of Certain Marine Organisms of (a) Variations in Reaction to Light, and (b) a Diurnal Periodicity of Phosphorescence. *Biochem. J.*, 1909, iv, 1-29.
- 371a MORGAN, C. L.: Animal Behavior. London, 1900.
- 371b MORGULIS, S.: The Auditory Reactions of the Dog Studied by the Pawlow Method. *J. Animal Behav.*, 1914, iv, 142-145.
- 371c MORGULIS, S.: Pawlow's Theory of the Function of the Central Nervous System and a Digest of Some of the More Recent Contributions to This Subject from Pawlow's Laboratory. *J. Animal Behav.*, 1914, iv, 362-379.
- 372 MORSE, M. W.: Alleged Rhythm in Phototaxis Synchronous with Ocean Tides. *Proc. Soc. Exp. Biol. and Med.*, 1910, vii, 145-146.
- 373 MÜLLER, H.: Ueber Heliotropismus. *Flora*, 1876, lix, 65-70, 88-95.
- 374 MÜLLER-HETTLINGEN, J.: Ueber galvanische Erscheinungen an keimenden Samen. *Arch. ges. Physiol.*, 1883, xxxi, 193-212.
- 375 MURBACH, L.: The Static Function in *Gonionemus*. *Am. J. Physiol.*, 1903, x, 201-209.
- 376 MURBACH, L.: Some Light Reactions of the Medusa *Gonionemus*. *Biol. Bull.*, 1909, xvii, 354-368.
- 377 MUSSET, CH.: Sélénotropisme. *Compt. rend. Acad. Sc.*, 1890, cx, 201-202.
- 378 NAGEL, W. A.: Beobachtungen über den Lichtsinn augenloser Muscheln. *Biol. Centr.*, 1894, xiv, 385-390.
- 379 NAGEL, W. A.: Ein Beitrag zur Kenntnis des Lichtsinnes augenloser Tiere. *Biol. Centr.*, 1894, xiv, 810-813.
- 379a NAGEL, W. A.: Experimentelle sinnesphysiologische Untersuchungen an Coelenteraten. *Arch. ges. Physiol.*, 1894, lvii, 495-552.
- 380 NAGEL, W. A.: Ueber Galvanotaxis. *Arch. ges. Physiol.*, 1895, lix, 603-642.
- 381 NAGEL, W. A.: Der Lichtsinn augenloser Tiere. Jena, 1896, pp. 120.
- 382 NAGEL, W. A.: Phototaxis, Photokinesis und Unterschiedsempfindlichkeit. *Bot. Ztg.*, 1901, lix, 298-299.

- 383 NAGEL, W. A.: Methoden zur Erforschung des Licht- und Farbensinnes. *Tigerstedt's Handb. physiol. Methodik*, 1909, iii, Abt. 2, *Sinnesphysiologie*, ii, 1-99.
- 384 NATHANSOHN, A., and PRINGSHEIM, E.: Ueber die Summation intermittierender Lichtreize. *Jahrb. wiss. Bot.*, 1908, xlv, 137-190.
- 385 NĚMEC, B.: Ueber die Wahrnehmung des Schwerkraftreizes bei den Pflanzen. *Jahrb. wiss. Bot.*, 1901, xxxvi, 80-178.
- 385a NERNST, W., and BARRATT, J. O. W.: Ueber die elektrische Nervenreizung durch Wechselströme. *Z. Electrochem.*, 1904, x, 664-668.
- 386 NEUBERG, C.: Chemische Umwandlungen durch Strahlenarten. *Biochem. Z.*, 1908, xiii, 305-320; 1909, xvii, 270-292.
- 387 NUEL, J. P.: *La vision*. Paris, 1904, pp. 376.
- 388 NYBERGH, T.: Studien über die Einwirkung der Temperatur auf die tropistische Reisbarkeit etiolierter *Avena*-Keimlinge. *Ber. bot. Ges.*, 1912, xxx, 542-553.
- 389 OLTMANNS, F.: Ueber die photometrischen Bewegungen der Pflanzen. *Flora*, 1892, lxxv, 183-266.
- 390 OLTMANNS, F.: Ueber positiven und negativen Heliotropismus. *Flora*, 1897, lxxxiii, 1.
- 391 OSTWALD, Wo.: Ueber eine neue theoretische Betrachtungsweise in der Planktologie, insbesondere über die Bedeutung des Begriffs der "inneren Reibung des Wassers" für dieselbe. *Forsch.-ber. biol. Station Ploën*, 1903, pt. 10, 1-49.
- 392 OSTWALD, Wo.: Zur Theorie der Richtungsbewegungen schwimmender niederer Organismen. *Arch. ges. Physiol.*, 1903, xcv, 23-65; 1906, cxi, 452-472; 1907, cxvii, 384-408.
- 393 OSTWALD, Wo.: Ueber die Lichtempfindlichkeit tierischer Oxydasen und über die Beziehungen dieser Eigenschaft zu den Erscheinungen des tierischen Phototropismus. *Biochem. Z.*, 1908, x, 1-130.
- 394 PAAL, A.: Ueber phototropische Reizleitungen. *Ber. bot. Ges.*, 1914, xxxii, 499-502.
- 395 PARKER, G. H.: Photomechanical Changes in the Retinal Pigment Cells of *Palæmonetes*, and Their Relation to the Central Nervous System. *Bull. Mus. Comp. Zool.*, 1897, xxx, 273-300.
- 396 PARKER, G. H.: The Photomechanical Changes in the Retinal Pigment of *Gammarus*. *Bull. Mus. Comp. Zool.*, 1899, xxxv, 141-148.
- 397 PARKER, G. H.: The Reactions of Copepods to Various Stimuli and the Bearing of This on Daily Depth-migrations. *Bull. U. S. Fish Comm.*, 1901, 103-123.
- 398 PARKER, G. H.: The Phototropism of the Mourning-cloak Butterfly, *Vanessa antiopa* Linn. *Mark Anniversary Vol.*, 1903, 453-469.

- 399 PARKER, G. H.: The Skin and the Eyes as Receptive Organs in the Reactions of Frogs to Light. *Am. J. Physiol.*, 1903, x, 28-36.
- 400 PARKER, G. H.: The Stimulation of the Integumentary Nerves of Fishes by Light. *Am. J. Physiol.*, 1905, xiv, 413-420.
- 401 PARKER, G. H.: The Reactions of *Amphioxus* to Light. *Proc. Soc. Exp. Biol. and Med.*, 1906, iii, 61-62.
- 402 PARKER, G. H.: The Influence of Light and Heat on the Movement of the Melanophore Pigment, Especially in Lizards. *J. Exp. Zool.*, 1906, iii, 401-414.
- 403 PARKER, G. H.: The Sensory Reactions of *Amphioxus*. *Proc. Am. Acad. Arts and Sc.*, 1908, xliii, 415-455.
- 404 PARKER, G. H.: The Integumentary Nerves of Fishes as Photoreceptors and Their Significance for the Origin of the Vertebrate Eyes. *Am. J. Physiol.*, 1909, xxv, 77-80.
- 405 PARKER, G. H.: Mast's "Light and the Behavior of Organisms." *J. Animal Behav.*, 1911, i, 461-464.
- 406 PARKER, G. H., and ARKIN, L.: The Directive Influence of Light on the Earthworm *Allolobophora fatida* (Sav.). *Am. J. Physiol.*, 1901, v, 151-157.
- 407 PARKER, G. H., and BURNETT, F. L.: The Reactions of *Planarians*, With and Without Eyes, to Light. *Am. J. Physiol.*, 1900, iv, 373-385.
- 408 PARKER, G. H., and METCALF, C. R.: The Reactions of Earthworms to Salts: a Study in Protoplasmic Stimulation as a Basis of Interpreting the Sense of Taste. *Am. J. Physiol.*, 1906, xvii, 55-74.
- 409 PARKER, G. H., and PARSHLEY, H. M.: The Reactions of Earthworms to Dry and to Moist Surfaces. *J. Exp. Zool.*, 1911, xi, 361-363.
- 410 PARKER, G. H., and PATTEN, B. M.: The Physiological Effect of Intermittent and of Continuous Lights of Equal Intensities. *Am. J. Physiol.*, 1912, xxxi, 22-29.
- 411 PARMLLEE, M.: The Science of Human Behavior. New York, 1913, xvii+443.
- 412 PATTEN, B. M.: A Quantitative Determination of the Orienting Reaction of the Blowfly Larva (*Calliphora erythrocephala* Meigen), *J. Exp. Zool.*, 1914, xvii, 213-280.
- 413 PATTEN, B. M.: An Analysis of Certain Photic Reactions with Reference to the Weber-Fechner Law. I. The Reactions of the Blowfly Larva to Opposed Beams of Light. *Am. J. Physiol.*, 1915, xxxviii, 313-338.
- 414 PATTEN, B. M.: The Changes of the Blowfly Larva's Photosensitivity with Age. *J. Exp. Zool.*, 1916, xx, 585-598.

- 415 PATTEN, B. M.: Reactions of the Whip-tail Scorpion to Light. *J. Exp. Zool.*, 1917, xxiii, 251-275.
- 416 PAYNE, F.: The Reactions of the Blind Fish, *Amblyopsis spelæus*, to Light. *Biol. Bull.*, 1907, xiii, 317-323.
- 416^a PAYNE, F.: Forty-nine Generations in the Dark. *Biol. Bull.*, 1910, xviii, 188-190.
- 416^b PAYNE, F.: *Drosophila ampelophila* Loew Bred in the Dark for Sixty-nine Generations. *Biol. Bull.*, 1911, xxi, 297-301.
- 417 PEARL, R.: Studies on Electrotaxis. I. On the Reactions of Certain Infusoria to the Electric Current. *Am. J. Physiol.*, 1900, iv, 96-123.
- 418 PEARL, R.: Studies on the Effects of Electricity on Organisms. II. The Reactions of *Hydra* to the Constant Current. *Am. J. Physiol.*, 1901, v, 301-320.
- 419 PEARL, R.: The Movements and Reactions of Fresh-water *Planarians*: a Study in Animal Behavior. *Quart. J. Micr. Sc.*, 1902-03, xlv, 509-714.
- 420 PEARL, R., and COLE, L. J.: The Effect of Very Intense Light on Organisms. *Third Rep. Mich. Acad. Sc.*, 1901, 77-78.
- 421 PEARSE, A. S.: The Reactions of Amphibians to Light. *Proc. Am. Acad. Arts and Sc.*, 1910, xlv, 161-208.
- 422 PÉREZ, J.: Notes zoologiques. De l'attraction exercée par les odeurs et les couleurs sur les insectes. *Acta Soc. Linn., Bordeaux*, 1894, vii, 245-253.
- 423 PFEFFER, W.: Locomotorische Richtungsbewegungen durch chemische Reize. *Ber. bot. Ges.*, 1883, i, 524-533.
- 424 PFEFFER, W.: Locomotorische Richtungsbewegungen durch chemische Reize. *Unters. Bot. Inst. Tübingen*, 1884, i, 363-482.
- 425 PFEFFER, W.: Ueber chemotaktische Bewegungen von Bakterien, Flagellaten und Volvocineen. *Unters. Bot. Inst. Tübingen*, 1888, ii, 582-661.
- 426 PHIPPS, C. F.: An Experimental Study of the Behavior of Amphipods with Respect to Light Intensity, Direction of Rays, and Metabolism. *Biol. Bull.*, 1915, xxviii, 210-223.
- 427 PLATEAU, F.: Recherches sur la perception de la lumière par les myriopodes aveugles. *J. Anat. et Physiol.*, 1886, xxii.
- 428 PLATEAU, F.: Nouvelles recherches sur les rapports entre les insectes et les fleurs. *Mém. Soc. Zool. France*, 1899, xii.
- 429 PLATEAU, F.: La choix des couleurs par les insectes. *Mém. Soc. Zool. France*, 1899, xii, 336-370.
- 430 PLATEAU, F.: Expériences sur l'attraction des insectes par les étoffes colorées et les objets brillants. *Ann. Soc. Ent. Belgique*, 1900, xlv.

- ⁴³¹ PLATT, J. B.: On the Specific Gravity of *Spirostomum*, *Paramæcium*, and the Tadpole in Relation to the Problem of Geotaxis. *Am. Nat.*, 1899, xxxiii, 31-38.
- ⁴³² POLIMANTI, O.: Ueber eine beim Phototropismus des *Lasius niger* L. beobachtete Eigentümlichkeit. *Biol. Centr.*, 1911, xxxi, 222-224.
- ⁴³³ POLIMANTI, O.: Sul reotropismo nelle larve dei batraci (*Bufo e Rana*). *Biol. Centr.*, 1915, xxxv, 36-39.
- ⁴³⁴ PORODKO, TH. M.: Vergleichende Untersuchungen über die Tropismen. I. Das Wesen der chemotropen Erregung bei den Pflanzenwurzeln. *Ber. bot. Ges.*, 1912, xxx, 16-27.
- ⁴³⁵ PORODKO, TH. M.: II. Thermotropismus der Pflanzenwurzeln. *Ber. bot. Ges.*, 1912, xxx, 305-313.
- ⁴³⁶ PORODKO, TH. M.: IV. Die Gültigkeit des Energiemengengesetzes für den negativen Chemotropismus der Pflanzenwurzeln. *Ber. bot. Ges.*, 1913, xxxi, 88-94.
- ⁴³⁷ PORODKO, TH. M.: V. Das mikroskopische Aussehen der tropistisch gereizten Pflanzenwurzeln. *Ber. bot. Ges.*, 1913, xxxi, 248-256.
- ⁴³⁸ POWERS, E. B.: The Reactions of Crayfishes to Gradients of Dissolved Carbon Dioxide and Acetic and Hydrochloric Acids. *Biol. Bull.*, 1914, xxvii, 177-200.
- ⁴³⁹ PRENTISS, C. W.: The Otocyst of Decapod Crustacea: Its Structure, Development, and Functions. *Bull. Mus. Comp. Zool.*, 1901, xxxvi, 165-251.
- ⁴⁴⁰ PRINGSHEIM, E. G.: Die Reizbewegungen der Pflanzen. Berlin, 1912, viii+326.
- ⁴⁴¹ PRINGSHEIM, E. G.: Das Zustandekommen der taktischen Reaktionen. *Biol. Centr.*, 1912, xxxii, 337-365.
- ⁴⁴² PRZIBRAM, K.: Ueber die ungeordnete Bewegung niederer Tiere. *Arch. ges. Physiol.*, 1913, cliii, 401-405.
- ⁴⁴³ PÜTTER, A.: Studien über Thigmotaxis bei Protisten. *Arch. Anat. u. Physiol.*, 1900, *Suppl.*, 243-302.
- ⁴⁴⁴ RÄDL, E.: Ueber den Phototropismus einiger Arthropoden. *Biol. Centr.*, 1901, xxi, 75-86.
- ⁴⁴⁵ RÄDL, E.: Untersuchungen über die Lichtreaktion der Arthropoden. *Arch. ges. Physiol.*, 1901, lxxvii, 418-466.
- ⁴⁴⁶ RÄDL, E.: Ueber die Lichtreaktionen der Arthropoden auf der Drehscheibe. *Biol. Centr.*, 1902, xxii, 728-732.
- ⁴⁴⁷ RÄDL, E.: Untersuchungen über den Phototropismus der Tiere. Leipzig, 1903, viii+188.
- ⁴⁴⁸ RÄDL, E.: Ueber die Anziehung des Organismus durch das Licht. *Flora*, 1904, xciii, 167-178.
- ⁴⁴⁹ RÄDL, E.: Einige Bemerkungen und Beobachtungen über den Phototropismus der Tiere. *Biol. Centr.*, 1906, xxvi, 677-690.

- 450 RÉAUMUR: Mémoires pour servir a l'histoire des insectes. Paris, 1740.
- 451 REESE, A. M.: Observations on the Reactions of *Cryptobranchus* and *Necturus* to Light and Heat. *Biol. Bull.*, 1906, xi, 93-99.
- 452 RILEY, C. F. C.: Observations on the Ecology of Dragon-fly Nymphs: Reactions to Light and Contact. *Ann. Ent. Soc. Am.*, 1912, v, 273-292.
- 453 ROMANES, G. J.: Animal Intelligence. New York, 1883, pp. 520.
- 454 ROMANES, G. J.: Jelly-fish, Star-fish and Sea-urchins. New York, 1893, x+323.
- 455 ROTHERT, W.: Ueber Heliotropismus. *Beitr. Biol. Pflanzen*, 1894, vii, 1.
- 456 ROTHERT, W.: Beobachtungen und Betrachtungen über taktische Reizerscheinungen. *Flora*, 1901, lxxxviii, 371-421.
- 457 ROUX, W.: Ueber die Selbstordnung (Cytotaxis) sich "berührender" Furchungszellen des Froscheies durch Zusammenfügung, Zellentrennung und Zellengleiten. *Arch. Entwicklungsmech.*, 1896, iii, 381-468.
- 458 ROYCE, J.: Outlines of Psychology. New York, 1903, pp. 417.
- 459 RUCHLÁDEW, N.: Untersuchungen zur Kritik der Methodik chemotaktischer Versuche und zur Biologie der Leukozyten. *Z. Biol.*, 1910, liv, 533-559.
- 460 SCHÄFER, K. L.: Ueber den Drehschwindel bei den Tieren. *Z. Psychol. u. Physiol. Sinnesorg.*, 1891.
- 461 SCHAEFFER, A. A.: Reactions of *Ameba* to Light and the Effect of Light on Feeding. *Biol. Bull.*, 1917, xxxii, 45-74.
- 462 SCHMID, B.: Ueber den Heliotropismus von *Cereactis aurantiaca*. *Biol. Bull.*, 1911, xxxi, 538-539.
- 462a SCHNEIDER, G. H.: Der tierische Wille. Leipzig, 1880.
- 462b SCHNEIDER, K. C.: Tierpsychologisches Praktikum in Dialogform. Leipzig, 1912, pp. 719.
- 462c SCHNEIDER, K. C.: Vorlesungen über Tierpsychologie. Leipzig, 1909.
- 463 SCHOENICHEN, W.: Die Empfindlichkeit der Nachtschmetterlinge gegen Lichtstrahlen. *Prometheus*, 1904, xvi, 29-30.
- 464 SCHOUTEDEN, H.: Le phototropisme de *Daphnia magna* Straus (Crust.). *Ann. Soc. Ent. Belgique*, 1902, xlii, 352-362.
- 465 SHIBATA, K.: Studien über die Chemotaxis der *Isoëtes*-Spermatozoiden. *Jahrb. wiss. Bot.*, 1905, xli, 561-610.
- 466 SHOHL, A. T.: Reactions of Earthworms to Hydroxyl Ions. *Am. J. Physiol.*, 1914, xxxiv, 384-404.

- ⁴⁶⁷ SMITH, A. C.: The Influence of Temperature, Odors, Light, and Contact on the Movements of the Earthworm. *Am. J. Physiol.*, 1902, vi, 459-486.
- ⁴⁶⁸ SMITH, G.: The Effect of Pigment-migration on the Phototropism of *Gammarus annulatus* S. I. Smith. *Am. J. Physiol.*, 1905, xiii, 205-216.
- ⁴⁶⁹ SOSNOWSKI, J.: Untersuchungen über die Veränderungen des Geotropismus bei *Paramæcium aurelia*. *Bull. Internat. Acad. Sc. Cracovie*, 1899, 130-136.
- ⁴⁷⁰ STATKEWITSCH, P.: Ueber die Wirkung der Induktionsschläge auf einige Ciliata. *Le Physiologiste Russe*, 1903, iii, 41-45.
- ⁴⁷¹ STATKEWITSCH, P.: Galvanotropismus und Galvanotaxis der Ciliata. *Z. allg. Physiol.*, 1904, iv, 296-332; 1905, v, 511-534; 1907, vi, 13-43.
- ⁴⁷² STRASBURGER, E.: Wirkung des Lichtes und der Wärme auf Schwärmsporen. *Jenaische Z. Naturwiss.*, 1878, (N.F.) xii, 551-625. Also separate, Jena, pp. 75.
- ⁴⁷³ SZYMANSKI, J. S.: Ein Versuch, das Verhältnis zwischen modal verschiedenen Reizen in Zahlen auszudrücken. *Arch. ges. Physiol.*, 1911, cxxxviii, 457-486.
- ⁴⁷⁴ SZYMANSKI, J. S.: Aenderung des Phototropismus bei Kiechenschaben durch Erlernung. *Arch. ges. Physiol.*, 1912, cxliv, 132-134.
- ⁴⁷⁵ SZYMANSKI, J. S.: Ein Beitrag zur Frage über tropische Fortbewegung. *Arch. ges. Physiol.*, 1913, cliv, 343-363.
- ⁴⁷⁶ SZYMANSKI, J. S.: Methodisches zum Erforschen der Instinkte. *Biol. Centr.*, 1913, xxxiii, 260-264.
- ⁴⁷⁷ v. TAPPEINER, H.: Die photodynamische Erscheinung (Sensibilisierung durch fluoreszierende Stoffe). *Ergeb. Physiol.*, 1909, viii, 698-741.
- ⁴⁷⁸ TERRY, O. P.: Galvanotropism of *Volvox*. *Am. J. Physiol.*, 1906, xv, 235-243.
- ⁴⁷⁹ TORELLI, E.: The Response of the Frog to Light. *Am. J. Physiol.*, 1903, ix, 466-488.
- ⁴⁸⁰ TORREY, H. B.: On the Habits and Reactions of *Sagartia davisi*. *Biol. Bull.*, 1904, vi, 203-216.
- ⁴⁸¹ TORREY, H. B.: The Method of Trial and the Tropism Hypothesis. *Science*, 1907, xxvi, 313-323.
- ⁴⁸² TORREY, H. B.: Trials and Tropisms. *Science*, 1913, xxxvii, 873-876.
- ⁴⁸³ TORREY, H. B.: Tropisms and Instinctive Activities. *Psychol. Bull.*, 1916, xiii, 297-308.
- ⁴⁸⁴ TORREY, H. B., and HAYS, G. P.: The Rôle of Random Movements in the Orientation of *Porcellio scaber* to Light. *J. Animal Behav.*, 1914, iv, 110-120.

- 485 TOWLE, E. W.: A Study in the Heliotropism of *Cypridopsis*. *Am. J. Physiol.*, 1900, iii, 345-365.
- 486 TURNER, C. H.: An Experimental Investigation of an Apparent Reversal of the Responses to Light of the Roach (*Periplaneta orientalis* L.). *Biol. Bull.*, 1912, xxiii, 371-386.
- 487 v. UEXKÜLL, J.: Vergleichend-sinnesphysiologische Untersuchungen. II. Der Schatten als Reiz für *Centrostephanus longispinus*. *Z. Biol.*, 1897, xxxiv, 319-339.
- 488 v. UEXKÜLL, J.: Die Wirkung von Licht und Schatten auf die Seeigel. *Z. Biol.*, 1900, xl, 447-476.
- 489 v. UEXKÜLL, J.: Umwelt und Innenwelt der Tiere. Berlin, 1909, pp. 261.
- 490 ULEHLA, VL.: Ultramikroskopische Studien über Geisselbewegung. *Biol. Centr.*, 1911, xxxi, 645-654, 657-676, 689-705, 721-731.
- 491 VAN HERWERDEN, M. A.: Ueber die Perzeptionsfähigkeit des Daphnienauges für ultra-violette Strahlen. *Biol. Centr.*, 1914, xxxiv, 213-216.
- 492 VERWORN, M.: Psycho-physiologische Protistenstudien. Experimentelle Untersuchungen. Jena, 1889, viii+219.
- 493 VERWORN, M.: Die polare Erregung der Protisten durch den galvanischen Strom. *Arch. ges. Physiol.*, 1889, xlv, 1-36; 1890, xlv, 267-303.
- 494 VERWORN, M.: Gleichgewicht und Otolithenorgan. Experimentelle Untersuchungen. *Arch. ges. Physiol.*, 1891, l, 423-472.
- 495 VERWORN, M.: Untersuchungen über die polare Erregung der lebendigen Substanz durch den konstanten Strom. *Arch. ges. Physiol.*, 1896, lxii, 415-450.
- 496 VERWORN, M.: Die polare Erregung der lebendigen Substanz durch den konstanten Strom. *Arch. ges. Physiol.*, 1896, lxv, 47-62.
- 497 VERWORN, M.: General Physiology. New York, 1899.
- 498 VIEWEGER, TH.: Recherches sur la sensibilité des infusoires (alcalio-oxytaxisme), les réflexes locomoteurs, l'action des sels. *Arch. Biol.*, 1912, xxvii, 723-799.
- 499 DE VRIES, H.: Ueber einige Ursachen der Richtung bilateralsymmetrischer Pflanzenteile. *Arb. bot. Inst. Würzburg*, 1872, i, 223.
- 500 DE VRIES, M. S.: Die phototropische Empfindlichkeit des Segerhafers bei extremen Temperaturen. *Ber. bot. Ges.*, 1913, xxxi, 233-237.
- 501 WAGER, H.: On the Effect of Gravity upon the Movements and Aggregation of *Euglena viridis* Ehrb., and Other Microorganisms. *Phil. Trans. Roy. Soc. London*, 1911, cci, (B), 333-390.
- 502 WALLENGREN, H.: Zur Kenntnis der Galvanotaxis. I. Die anodische Galvanotaxis. *Z. allg. Physiol.*, 1903, ii, 341-384.

- 503 WALLENGREN, H.: II. Eine Analyse der Galvanotaxis bei *Spirostomum*. *Z. allg. Physiol.*, 1903, ii, 516-555.
- 504 WALLENGREN, H.: III. Die Entwirkung des konstanten Stromes auf die inneren Protoplasmabewegungen bei den Protozoen. *Z. allg. Physiol.*, 1904, iii, 22-32.
- 505 WALTER, H. E.: The Reactions of *Planarians* to Light. *J. Exp. Zool.*, 1907, v, 35-162.
- 506 WASHBURN, M. F.: The Animal Mind. New York, 1909, pp. 333.
- 507 WEIGERT, F.: Die chemischen Wirkungen des Lichts. Stuttgart, 1911.
- 508 WHEELER, W. M.: Anemotropism and Other Tropisms in Insects. *Arch. Entwicklungsmech.*, 1899, viii, 373-381.
- 509 WHITMAN, C. O.: Animal Behavior. Woods Hole Biol. Lectures, Boston, 1899, 285-338.
- 510 DE WILDEMAN, E.: Sur le thermotaxisme des *Euglènes*. *Bull. Soc. Belg. Micros.*, 1894, xx, 245-258.
- 511 V. WIESNER, J.: Heliotropismus und Strahlengang. *Ber. bot. Ges.*, 1912, xxx, 235-245.
- 512 WILLEM, V.: La vision chez les gastropodes pulmonés. *Compt. rend. Acad. Sc.*, 1891, cxii, 247-248.
- 513 WILLEM, V.: Sur les perceptions dermatoptiques. *Bull. Sc. France et Belgique*, 1891, xxiii, 329-346.
- 514 WILSON, E. B.: The Heliotropism of *Hydra*. *Am. Nat.*, 1891, xxv, 413-433.
- 515 WODSEDALEK, J. E.: Phototactic Reactions and Their Reversal in the May-fly Nymphs *Heptagenia interpunctata* (Say.). *Biol. Bull.*, 1911, xxi, 265-271.
- 516 YERKES, R. M.: Reaction of Entomostraca to Stimulation by Light. I. *Am. J. Physiol.*, 1899, iii, 157-182.
- 517 YERKES, R. M.: II. Reactions of *Daphnia* and *Cypris*. *Am. J. Physiol.*, 1900, iv, 405-422.
- 518 YERKES, R. M.: A Study of the Reactions and the Reaction Time of the Medusa *Gonionemus murbachii* to Photic Stimuli. *Am. J. Physiol.*, 1903, ix, 279-307.
- 519 YERKES, R. M.: Reactions of *Daphnia pulex* to Light and Heat. *Mark Anniversary Vol.*, 1903, 361-377.
- 520 YERKES, R. M.: The Reaction Time of *Gonionemus murbachii* to Electric and Photic Stimuli. *Biol. Bull.*, 1904, vi, 84-95.
- 521 ZAGOROWSKI, P.: Die Thermotaxis der *Paramécien*. *Z. Biol.*, 1914, lxxv, 1-12.
- 522 ZELIONY, G. P.: Observations sur des chiens auxquels on a enlevé les hémisphères cérébraux. *Compt. rend. Soc. Biol.*, 1913, lxxiv, 707-708.

- 523 BLASIUS, E., and SCHWEIZER, F.: Elektrotropismus und verwandte Erscheinungen. *Arch. ges. Physiol.*, 1893, liii, 493-543.
- 524 NERNST, W., and BARRATT, J. O. W.: Ueber die elektrische Nervenreizung durch Wechselströme. *Z. Electrochem.*, 1904, x, 664-668.
- 525 MOORE, A. R.: The Action of Strychnine on Certain Invertebrates. *J. Pharm. and Exp. Therap.*, 1916, ix, 167-169.
- 526 LOEB, J.: The Chemical Basis of Regeneration and Geotropism. *Science*, 1917, xlii, 115-118.
- 527 BREUER, J.: Ueber den Galvanotropismus (Galvanotaxis bei Fischen. *Sitzungsber. Akad. Wiss. Wien, mathem.-naturw. Kl.*, 1905, cxiv, 27-56.
- 528 BREUER, J., and KREIDL, A.: Ueber die scheinbare Drehung des Gesichtsfeldes, während der Einwirkung einer Centrifugalkraft. *Arch. ges. Physiol.*, 1898, lxx, 494-510.
- 529 HERMANN, L., and MATTHIAS, F.: Der Galvanotropismus der Larven von *Rana temporaria* und der Fische. *Arch. ges. Physiol.*, 1894, lvii, 391-405.
- 530 JENSEN, P.: Ueber den Geotropismus niederer Organismen. *Arch. ges. Physiol.*, 1893, liii, 428-480.
- 531 CROZIER, W. J.: The Photic Sensitivity of *Balanoglossus*. *J. Exp. Zool.*, 1917, xxiv, 211-217.
- 532 CLAPARÈDE, E.: Les tropismes devant la psychologie. *J. Psychol. u. Neurol.*, 1908, xiii, 150-160.
- 533 NAGEL, W. A.: Experimentelle sinnesphysiologische Untersuchungen an Coelenteraten. *Arch. ges. Physiol.*, 1894, lvii, 495-552.
- 534 SCHNEIDER, G. H.: Der tierische Wille. Leipzig, 1880.
- 535 SCHNEIDER, K. C.: Tierpsychologisches Praktikum in Dialogform. Leipzig, 1912, pp. 719.
- 536 SCHNEIDER, K. C.: Vorlesungen über Tierpsychologie. Leipzig, 1909.
- 537 MORGULIS, S.: The Auditory Reactions of the Dog Studied by the Pawlow Method. *J. Animal Behav.*, 1914, iv, 142-145.
- 538 MORGULIS, S.: Pawlow's Theory of the Function of the Central Nervous System and a Digest of Some of the More Recent Contributions to This Subject from Pawlow's Laboratory. *J. Animal Behav.*, 1914, iv, 362-379.
- 539 CRAIG, W.: The Voices of Pigeons Regarded as a Means of Social Control. *Am. J. Sociology*, 1908, xiv, 86-100.
- 540 CRAIG, W.: Male Doves Reared in Isolation. *J. Animal Behav.*, 1914, iv, 121-133.
- 541 MATULA, J.: Untersuchungen über die Funktionen des Zentralnervensystems bei Insekten. *Arch. ges. Physiol.*, 1911, cxxxviii, 388-456.

- ⁵⁴² LOEB, J.: Influence of the Leaf upon Root Formation and Geotropic Curvature in the Stem of *Bryophyllum calycinum* and the Possibility of a Hormone Theory of These Processes. *Bot. Gaz.*, 1917, lxiii, 25-50.
- ⁵⁴³ LOEB, J.: Untersuchungen zur physiologischen Morphologie der Tiere. I. Heteromorphose. II. Organbildung und Wachstum. Würzburg, 1891-1892.
- ⁵⁴⁴ LOEB, J.: The Chemical Mechanism of Regeneration. *Ann. Inst. Pasteur*, 1918, xxxii, 1-16.
- ⁵⁴⁵ CRAIG, W.: Appetites and Aversions as Constituents of Instincts. *Biol. Bull.*, 1918, xxxiv, 91-107.
- ⁵⁴⁶ KANDA, S.: Further Studies on the Geotropism of *Paramæcium caudatum*. *Biol. Bull.*, 1918, xxxiv, 108-119.
- ⁵⁴⁷ LYON, E. P.: Note on the Geotropism of *Paramæcium*. *Biol. Bull.*, 1918, xxxiv, 120.
- ⁵⁴⁸ McCLENDON, J. F.: Protozoan Studies. *J. Exp. Zool.*, 1909, vi, 265-283.
- ⁵⁴⁹ McEWEN, R. S.: The Reactions to Light and to Gravity in *Drosophila* and its Mutants. *J. Exp. Zool.*, 1918, xxv, 49-106.
- ⁵⁵⁰ PAYNE, F.: Forty-nine Generations in the Dark. *Biol. Bull.*, 1910, xviii, 188-190.
- ⁵⁵¹ PAYNE, F.: *Drosophila ampelophila* Loew Bred in the Dark for Sixty-nine Generations. *Biol. Bull.*, 1911, xxi, 297-301.
- ⁵⁵² MORGAN, C. L.: Animal Behavior. London, 1900.
- ⁵⁵³ STEVENS, N. M.: Regeneration in *Antennularia*. *Arch. Entwicklungs-mech.*, 1910, xxx, pt. 1, 1-7.
- ⁵⁵⁴ MAXWELL, S. S.: On the Exciting Cause of Compensatory Movements. *Am. J. Physiol.*, 1911-12, xxix, 367-371.

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